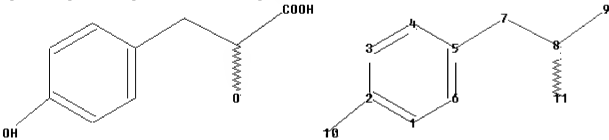
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ring nodes:
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ring bonds:
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exact bonds:
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normalized bonds:
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Match level:
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS 11:CLASS

L6 STRUCTURE UPLOADED

=> d 16
L6 HAS NO ANSWERS
L6 STR
/ Structure 1 in file .gra /

Structure attributes must be viewed using STN Express query preparation.

=> s 16 fam sam
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SAMPLE SCREEN SEARCH COMPLETED - 139 TO ITERATE

100.0% PROCESSED 139 ITERATIONS 1 ANSWERS SEARCH TIME: 00.00.01

SEARCH TIME. 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 2073 TO 3487
PROJECTED ANSWERS: 1 TO 80

=> s 16 fam full

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SEARCH TIME: 00.00.01

L8 16 SEA FAM FUL L6

=> s 16 sss sam

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SAMPLE SCREEN SEARCH COMPLETED - 597 TO ITERATE

100.0% PROCESSED 597 ITERATIONS 26 ANSWERS

16 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 10475 TO 13405 PROJECTED ANSWERS: 215 TO 825

L9 26 SEA SSS SAM L6

=> s 16 sss full

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SEARCH TIME: 00.00.01

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L14 26 SEA SSS SAM L12

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FULL SEARCH INITIATED 09:42:14 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 12087 TO ITERATE

100.0% PROCESSED 12087 ITERATIONS 412 ANSWERS

SEARCH TIME: 00.00.01

L15 412 SEA SSS FUL L12

=> d scan

L15 412 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN

IN D-Tyrosine, β -ethyl- α -[(4-fluorobenzoyl)oxy]-3-hydroxy-

MF C18 H18 F N O

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3034 L15

4705416 PREP/RL

L16 617 L15/PREP

(L15 (L) PREP/RL)

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L20 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                         2003:22888 CAPLUS Full-text
DOCUMENT NUMBER:
                         138:73085
TITLE:
                         Preparation of 3-aryl-\alpha-oxy substituted
                         propanoic acids for treatment of type II diabetes and
                         related disorders
                         Potlapally, Rajender Kumar; Velagala, Venkata Rama
INVENTOR(S):
                         Murali Krishna Reddy; Mamillapalli, Ramabhadra Sarma;
                         Gaddam, Om Reddy
PATENT ASSIGNEE(S):
                         Reddy's Research Foundation, India
                         PCT Int. Appl., 54 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Pat.ent.
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LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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                             DATE
                                                             DATE
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                                      WO 2001-IN124
    WO 2003002575
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                                                              20040728 <--
PRIORITY APPLN. INFO.:
                                        WO 2001-IN124
                                                          W 20010628 <--
OTHER SOURCE(S): CASREACT 138:73085; MARPAT 138:73085
GΙ
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/ Structure 28 in file .gra /

AB 3-Aryl- α -oxy substituted propanoic acids [I; wherein R1 = tertbutyldimethylsilyl, trimethylsilyl, alkoxyalkyl; R2 = H, (substituted) (C1-C6) alkyl] were prepared For example, Me tert-butyldimethylsilyloxy-3-(4-hydroxyphenyl) propanoate (II) was prepared in three steps. The prepared compds. are useful for treating diabetes, obesity, glucose intolerance, insulin resistance and other related disorders such as hypertension, coronary heart disease, atherosclerosis, stroke, peripheral vascular diseases and related disorders (no data). The prepared compds. are also useful for reducing total cholesterol, body weight, blood plasma glucose, triglycerides, LDL, VLDL and free fatty acids (no data).

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L20 ANSWER 2 OF 7 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:620035 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 138:89669

TITLE: Efficient synthesis of antihyperglycemic

(S)- α -aryloxy- β -phenylpropionic acid derivative using a bifunctional asymmetric

catalyst

AUTHOR(S): Takamura, Makoto; Yanagisawa, Hiroaki; Kanai, Motomu;

Shibasaki, Masakatsu

CORPORATE SOURCE: Medicinal Chemistry Research Laboratories, Sankyo Co.,

Ltd., Tokyo, 140-8710, Japan

SOURCE: Chemical & Pharmaceutical Bulletin (2002),

50(8), 1118-1121

CODEN: CPBTAL; ISSN: 0009-2363 Pharmaceutical Society of Japan

DOCUMENT TYPE: Journal LANGUAGE: English

PUBLISHER:

OTHER SOURCE(S): CASREACT 138:89669

/ Structure 29 in file .gra /

AB The title acid (I) was prepared using catalytic asym. cyanosilylation as a key reaction to construct the α -oxycarboxylic acid moiety.

REFERENCE COUNT: 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L20 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:286131 CAPLUS Full-text

DOCUMENT NUMBER: 136:309765

TITLE: Preparation of optically active 2-hydroxy-3-phenylpropionitrile

INVENTOR(S): Yanagisawa, Hiroaki; Takamura, Minoru

PATENT ASSIGNEE(S): Sankyo Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE ____ _____ _____ JP 2002114749 Α 20020416 JP 2000-303538 20001003 <--PRIORITY APPLN. INFO.: JP 2000-303538 20001003 <--

OTHER SOURCE(S): CASREACT 136:309765; MARPAT 136:309765

The compds. p-R10C6H4CH2CH(OH)CN (R1 = H, OH-protecting group) are prepared by reaction of p-R10C6H4CH2CHO (R1 = same as above) with cyanating agents in the presence of optically active catalysts. 4-Benzyloxyphenylacetaldehyde was reacted with trimethylsilyl cyanide in the presence of Et2AlCl, (S)-3,3'-bis(diphenylphosphinoyl)-1,1'- binaphthol, and Bu3P(O) in CH2Cl2 at -40° for 53.5 h to give 116 mg (R)-3-(4-benzyloxyphenyl)-2-hydroxypropionitrile. (R)-3-(4-hydroxyphenyl)lactic acid was prepared from the compound

L20 ANSWER 4 OF 7 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1999:784755 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 132:166053

TITLE: Total syntheses of all four stereoisomers of piscidic acid via catalytic asymmetric dihydroxylation of (Z)-

and (E)-trisubstituted olefins

AUTHOR(S): Toshima, Hiroaki; Saito, Masatoshi; Yoshihara,

Teruhiko

CORPORATE SOURCE: Department of Bioscience and Chemistry, Faculty of

Agriculture, Hokkaido University, Sapporo, 060-8589,

Japan

SOURCE: Bioscience, Biotechnology, and Biochemistry (

1999), 63(11), 1934-1941 CODEN: BBBIEJ; ISSN: 0916-8451

PUBLISHER: Japan Society for Bioscience, Biotechnology, and

Agrochemistry

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 132:166053

AΒ Four stereoisomers of (2S,3R)-(+)-piscidic acid were synthesized with high optical purity via Sharpless catalytic asym. dihydroxylation of (Z)- and (E)trisubstituted olefins in 6 steps from (4-hydroxyphenyl)pyruvic acid. That is Wittig reaction of Me (4-hydroxyphenyl)pyruvate with (carbomethoxymethylene)triphenylphosphorane give (Z)- and (E)-trisubstituted olefins in a 3:1 ratio, protecting the phenolic hydroxyl group with tertbutyldimethylsilyl, then the (Z)-olefin was subjected to asym. dihydroxylation by using the chiral ligand, dihydroquinidine 1,4-anthraquinonediyl diether, gave the product with 89% e.e. Desilylation and subsequent alkaline hydrolysis gave (2S, 3R)-(+)-piscidic acid with > 99% e.e. after recrystn. Use of ligand, dihydroquinine 1,4-anthraquinonediyl diether, gave (2R,3S)-(-)piscidic acid . Asym. dihydroxylation of the (E)-olefin with phthalazine ligands (dihydroquinidine and dihydroquinine 1,4-phthalazinediyl diethers) also gave high e.e. values product, followed by the same procedure mentioned above, gave (2S,3S)-(+)-3-epi-piscidic acid and (2R,3R)-(-)-2-epi-piscidic acid resp.

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L20 ANSWER 5 OF 7 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1996:262036 CAPLUS Full-text

DOCUMENT NUMBER: 124:289000

ORIGINAL REFERENCE NO.: 124:53583a,53586a

TITLE: Hydrogenation process and catalysts for the

production of phenyllactic acids from phenylpyruvic

acids

INVENTOR(S):
Morita, Hikari; Mori, Hiroyuki

PATENT ASSIGNEE(S): Nitto Chemical Industry Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 8 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
EP 696566	A1	19960214	EP 1995-112540		19950809 <
EP 696566 R: CH, DE, LI	B1	19980610			
JP 08053394	А	19960227	JP 1994-206102		19940809 <
JP 3394606	В2	20030407			
US 5684186	A	19971104	US 1995-511152		19950804 <
CN 1122325	A	19960515	CN 1995-109050		19950809 <
CN 1083422	С	20020424			
PRIORITY APPLN. INFO.:			JP 1994-206102	Α	19940809 <
OTHER SOURCE(S):	CASREA	CT 124:28900	00; MARPAT 124:289000		
GI					

/ Structure 30 in file .gra /

AB Phenyllactic acids [I; R1, R2 = H, OH, (un)branched alkyl, (un)branched alkoxy; R3 = H, (un)branched alkyl; R1R2 = methylenedioxy], useful as intermediates in the preparation of agrochems. (no data) and pharmaceuticals (no data), are prepared in high yield by the hydrogenation of phenylpyruvic acids (II) in the presence of a catalyst containing ≥1 Group VIII metal (e.g., Pd, Pt, Ni, etc.). Thus, 3-(4-hydroxyphenyl)pyruvic acid was hydrogenated in

MeOH using a Pd/C catalyst at $25^{\circ}/5$ kg/cm2, producing 3-(4-hydroxyphenyl)lactic acid.

L20 ANSWER 6 OF 7 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1993:603784 CAPLUS Full-text

DOCUMENT NUMBER: 119:203784

ORIGINAL REFERENCE NO.: 119:36369a,36372a

TITLE: Selectivity and specificity in substrate binding to

proteases: novel hydrolytic reactions catalyzed by α -chymotrypsin suspended in organic solvents

 $\alpha-\text{chymotrypsin}$ suspended in organic solvents with low water content and mediated by ammonium

hydrogen carbonate

AUTHOR(S): Ricca, Jean Marc; Crout, David H. G.

CORPORATE SOURCE: Dep. Chem., University of Warwick, Coventry, CV4 7AL,

UK

SOURCE: Journal of the Chemical Society, Perkin Transactions

1: Organic and Bio-Organic Chemistry (1972-1999) (

1993), (11), 1225-33

CODEN: JCPRB4; ISSN: 0300-922X

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 119:203784

AB α -Chymotrypsin suspended in organic solvents with low water content catalyzed hydrolytic reactions in the presence of ammonium hydrogen carbonate. Mol. modeling studies were carried out and structure-reactivity relationships were estimated by studying the hydrolysis of amino acid derivs. and analogs. The enzyme was stereoselective with respect to the hydrolysis of L-amino acid derivs., but no stereoselectivity was observed when α -hydroxy esters were used as substrates. A general procedure for the resolution of aromatic amino acid esters is given. The results are interpreted in terms of mol. modeling based on x-ray crystallog. data and literature data.

L20 ANSWER 7 OF 7 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1955:84086 CAPLUS Full-text

DOCUMENT NUMBER: 49:84086

ORIGINAL REFERENCE NO.: 49:15798a-i,15799a-c

TITLE: Constituents of Cortex piscidiae erythrinae. II.

Synthesis of O-methylpiscidic acid

AUTHOR(S): Buckle, A. L. J.; McGookin, Alexander; Robertson,

Alexander

CORPORATE SOURCE: Univ. Liverpool, UK

SOURCE: Journal of the Chemical Society (1954)

3981-6

CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

cf. C.A. 42, 4981f. Piscidic acid was shown previouslyto be (+)-p-HOC6H4CH2C(OH)(CO2H)CH(OH)CO2H (loc. cit.) and now has been confirmed by the synthesis of p-methoxybenzyltartaric acid, identical with natural O-Me piscidic acid. Several routes for the synthesis of tartaric acids of this type were examined Et α -phenylacetoacetate, b0.8 110° (2,4-dinitrophenylhydrazone, m. 94-5°), prepared by the method of Attwood, et al. (C.A. 17, 3183), with Pb(OAc)4 in HOAc, gave Et α -acetoxy- α -phenylacetoacetate, b0.2 128-30°. Similarly, from Et γ -phenylacetoacetate there resulted Et α -acetoxy- γ -phenylacetoacetate (I), b0.5 143-5°. I was obtained also from phenylacetyl bromide, Et diazoacetate, and HOAc. A mixture of I, HCN, and NaOH after 12 h. was diluted with EtOH, saturated with HCl,

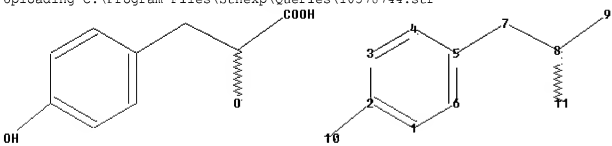
refluxed 4 h. and filtered to remove NH4Cl. The residue after evaporation of the filtrate gave Et benzyltartrate as a mixture which was separated manually into equal amts. of racemate A, m. $174-5^{\circ}$ (diamide, $204-6^{\circ}$), and racemate B, m. $194-5^{\circ}$ (diamide, m. $185-6^{\circ}$). Reduction of Et phenyloxaloacetate with moist Al amalgam gave Et β -phenylmalate, b0.04 131°, which after hydrolysis with KOH and purification from EtOAc gave β -phenylmalic acid, m. 150-60°. Fractional crystallization from EtOAc gave the racemic isomeride A, m. 172° , and from EtOAc-light petroleum (b.p. $60-80^{\circ}$) the racemic isomeride B, m. 162° . Dehydration with Ac20 of a mixture of the isomerides gave phenylmaleic anhydride, m. 120°, which on treatment with alkali gave phenylmaleic acid (II), m. $90-2^{\circ}$. II with pyridine and OsO4 in Et2O gave phenyltartaric acid, m. $173-4^{\circ}$. Oxidation of citraconic acid with NaClO3 and OsO4 gave Cmethyltartaric acid, m. $144-5^{\circ}$ (m. 146° , given by Schmidt and Perkow, C.A. 45, 2412h); Me ester, m. $99-100^{\circ}$; diamide, m. $152-3^{\circ}$. The condensation of NaOEt and Et oxalate with Et β -p-methoxyphenylpropionate, gave Et α -ethoxalyl- β -pmethoxyphenylpropionate. After reduction with moist amalgam, there was isolated Et β -hydroxy- α -p-methoxybenzylsuccinate, b0.1. 75-7°. Hydrolysis with KOH gave a mixture of acids, m. about 130°. Fractional crystallization from EtOAc-light petroleum (b.p. $60-80^{\circ}$) gave racemate A, m. $136-7^{\circ}$, and racemate B, m. $125-6^{\circ}$. A mixture of these racemates with Ac2O gave pmethoxybenzylidenesuccinic anhydride (III), $m. 160^{\circ}$, which on boiling with H2O gave p-methoxybenzylidenesuccinic acid (IV), m. and mixed m.p. 194-5° (decomposition). IV was obtained also from the condensation of anisaldehyde, Et succinate, and NaOEt. IV with Ac2O gave III, which by the boiling MeOH-H2SO4 method gave Me p-methoxybenzylidenesuccinate (V), b0.5 165°. Hydrogenation with PdC12 catalyst of IV gave p-methoxybenzylsuccinic acid (VI) m. 98-101°, and of V gave Me p-methoxybenzylsuccinate, bl 156°, m. 35-7°. Distillation of VI at $180^{\circ}/0.5$ mm. gave the anhydride, m. $91-2^{\circ}$. From pmethoxybenzyl alc. with PCl3 in Et20 there was obtained p-methoxybenzyl chloride as an unstable oil, $b25\ 125-7^{\circ}$, which on condensation with Et sodiomalonate gave Et p-methoxybenzylmalonate (VII), b0.5 145°. There was isolated from BrCH2CO2Et and VII Et α -ethoxycarbonyl- α -pmethoxybenzylsuccinate, b0.2 $166-9^{\circ}$ which on heating with EtOH-KOH gave α $carboxy-\alpha-p-methoxybenzylsuccinic acid, m. 157-9° (decomposition). When this$ was heated at $160^{\circ}/25$ mm. for 15 min., there was obtained VI. A stirred mixture of N-bromosuccinimide, p-methoxybenzylsuccinic anhydride, benzoyl peroxide, and CC14 or CS2 as solvent, after refluxing for 12 h., evaporating the filtered mixture and extracting with EtOAc gave III. III was heated until molten, then rapidly poured on to a cold surface, the solid pulverized and refluxed with CS2, collected and washed with more solvent and the process repeated 22 times. Evaporation of the combined CS2 exts. left an orange semisolid which was extracted with Et20. The residue left on evaporation of Et20 was extracted with light petroleum (b.p. $40-60^{\circ}$) and on cooling, deposited p-methoxybenzylmaleic anhydride (VIII), m. 64-5°, which on recrystn. from CHCl3-light petroleum (b.p. $60-80^{\circ}$), m. $65-6^{\circ}$. VIII reverted to III on melting. Hydrolysis of VIII with H2O gave p-methoxybenzylmaleic acid (IX), m. 120° (sintered at 117°). Addition of pyridine and OsO4 to IX in Et2O and the mixture kept in a closed vessel for 3 days resulted in a brown precipitate, which after collection was treated with aqueous KOH, the solution extracted with Et20, acidified, evaporated, the residue extracted with Et20 in a Soxhlet apparatus 9 h. and the extract evaporated to obtain p-methoxybenzyltartaric acid (X), m. 205-7° (decomposition); brucine salt, $[\alpha]23.5D$ -14.39° ± 0.6° (c 2.96, 50% EtOH). Resolution of X with brucine gave p-O-methylpiscidic acid, [α] 23D 44.01° \pm 5.0° (c 1.262, H2O), m. 169-70° (mixed m.p. with X, 173-6°); cinchonine salt, $[\alpha]17D$ 139.6° (c 6.1, EtOH); brucine salt, $[\alpha]24D$ -13.03° (c 2.131, 50% EtOH); Me ester, $[\alpha]$ 18D 78.16° (c 1.54, EtOH). The following derivs. of piscidic acid were cited: Me ester, $[\alpha]23D$ 41.52° (c 1.325, H2O); Et ester, $[\alpha]17.5D$ 59.70° (c 1.551, EtOH); di-Me ester, $[\alpha]19D$ 23.71° (c

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6.367, EtOH); Me p-O-benzylpiscidate, [\alpha]19D 48.73° (c 1.786, EtOH); cinchonine salt, [\alpha]21D 146.2° (c 0.424, EtOH); (+)-N-methylphenylisopropylamine salt, m. 179°, [\alpha]24D 12.73° (c 2.09, H2O). Reduction of p-methoxyphenylpyruvic acid in aqueous NaOH with 2% Na-Hg gave p-methoxyphenyllactic acid, m. 88°; Me ester (XI), b0.1 135°. Methylation of XI with Ag2O in MeI gave the Me ether of Me p-methoxyphenyllactate, b0.5 120°.
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http://www.cas.org/legal/infopolicy.html

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L32
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ring bonds :
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exact bonds :
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normalized bonds :
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11:CLASS
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                                                               412 ANSWERS
SEARCH TIME: 00.00.01
           412 SEA SSS FUL L42
L43
=> d scan
L43 412 ANSWERS
                  REGISTRY COPYRIGHT 2009 ACS on STN
TN
     Benzenepropanoic acid, 4-hydroxy-\alpha-methoxy-, (\alphaR)-(2S)-compd.
     with 2-amino-1-butanol (1:1)
    C10 H12 O4 . C4 H11 N O
    CM
         1
Absolute stereochemistry.
/ Structure 34 in file .gra /
=> s 143/prep
          3034 L43
       4705416 PREP/RL
           617 L43/PREP
L44
                 (L43 (L) PREP/RL)
=> s 144 and 'chiral catalyst'
        129174 'CHIRAL'
            19 'CHIRALS'
        129179 'CHIRAL'
                ('CHIRAL' OR 'CHIRALS')
        827404 'CATALYST'
        823540 'CATALYSTS'
       1060120 'CATALYST'
                 ('CATALYST' OR 'CATALYSTS')
          2138 'CHIRAL CATALYST'
                 ('CHIRAL'(W)'CATALYST')
L45
             0 L44 AND 'CHIRAL CATALYST'
=> s 144 and "asymmetric hydrogenation"
MISMATCHED QUOTE 'AND "ASYMMETRIC'
Quotation marks (or apostrophes) must be used in pairs,
one before and one after the expression you are setting
off or masking.
=> s 144 and 'asymmetric hydrogenation'
         76804 'ASYMMETRIC'
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31 'ASYMMETRICS'

76835 'ASYMMETRIC'

('ASYMMETRIC' OR 'ASYMMETRICS')

147804 'ASYM'

6 'ASYMS'

147807 'ASYM'

('ASYM' OR 'ASYMS')

171488 'ASYMMETRIC'

('ASYMMETRIC' OR 'ASYM')

184214 'HYDROGENATION'

2478 'HYDROGENATIONS'

184475 'HYDROGENATION'

('HYDROGENATION' OR 'HYDROGENATIONS')

4061 'ASYMMETRIC HYDROGENATION'

('ASYMMETRIC'(W)'HYDROGENATION')

L46 2 L44 AND 'ASYMMETRIC HYDROGENATION'

=> d 146 abs ibib 1-2

L46 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN

Chiral 2-alkoxy-3-arylpropanoic acids R2nC6H5-nCH2CH(OR1)CO2H or their alkali metal salts [1; R1 = (un)substituted C1-18 alkyl, C4-24 aryl, C5-18 arylalkyl; R2 = OH, halo, (alkyl)amino, C1-18 alkyl(oxy), C4-24 aryl, C5-18 arylalkyl, C1-18 alkylsulfonyl(amino), acyl(amino), acyloxy, preferably R2 = OH; n = 1-5, preferably n = 1, useful as peroxisome proliferator activated receptors (PPAR) agonists, were prepared by an improved process comprising transition metal-catalyzed asym. hydrogenation of the corresponding cinnamic acids R2nC6H5-nCH:C(OR1)CO2H (2; same R, n) in the presence of at least one protic solvent. Compds. 2 were preferably prepared by Perkin condensation of benzaldehydes R2nC6H5-nCHO (3; same R2, n) with 2-alkoxyacetates R1OCH2CO2R3 (4; same R1; R3 = H, C1-18 alkyl, preferably C1-6 alkyl). In an example, sodium 4-hydroxy- α -methoxybenzenepropanoate (α S)-4-HOC6H4CH2CH(OMe)CO2Na was prepared in 53% yield and 92% ee by asym. hydrogenation of 200.0 mmol of (2Z)-4-HOC6H4CH:C(OMe)CO2H catalyzed by 0.5 mmol of [Ir(COD)Cl]2 and 1.0 mmol of (S,S)-2,4-bis(diphenylphosphino)pentane in 240 mL of iso-Pr acetate and 60 mL of MeOH for 24 h at 65° and 3 atm of H2.

ACCESSION NUMBER: 2007:696717 CAPLUS Full-text

DOCUMENT NUMBER: 147:95305

TITLE: Process for the preparation of enantiomer-enriched

2-alkoxy-3-arylpropionic acids by asymmetric hydrogenation of substituted 2-alkoxycinnamic

acids

INVENTOR(S): Woltering, Michael; Bunlaksananusorn, Tanasri;

Gerlach, Arne

PATENT ASSIGNEE(S): Saltigo G.m.b.H., Germany SOURCE: Eur. Pat. Appl., 16pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
EP 1801093	A1 20070627	EP 2006-25546	20061211
R: AT, BE, BG,	CH, CY, CZ, DE,	DK, EE, ES, FI, FR, GB,	GR, HU, IE,
IS, IT, LI,	LT, LU, LV, MC,	NL, PL, PT, RO, SE, SI,	SK, TR, AL,
BA, HR, MK,	YU		
DE 102005061472	A1 20070705	DE 2005-102005061472	20051222
US 20070149804	A1 20070628	US 2006-635302	20061207
US 7429676	B2 20080930		

CN 1986516 A 20070627 CN 2006-10168677 20061222 PRIORITY APPLN. INFO.: DE 2005-102005061472A 20051222

OTHER SOURCE(S): CASREACT 147:95305; MARPAT 147:95305

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L46 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN GI

/ Structure 37 in file .gra /

Title compds. (I; R2 = alkyl; R5-R8 = H, substituent) and salts thereof were prepared by reaction of benzaldehydes (II; R1 = protective group; R5-R8 as defined above) with R2OCH2CO2R3 (R3 = hydrocarbyl; R2 as defined above), hydrolysis of the resulting cinnamate esters to give cinnamic acids, asym. hydrogenation, and O-deprotection. Thus, a mixture of 4-benzyloxybenzaldehyde, Me methoxyacetate, and NaOMe was refluxed 5 h in MeOH to give 80% Me 3-(4-benzyloxyphenyl)-2-methoxyacrylate. This was refluxed 2 h with 1N NaOH in MeOH to give 85% 3-(4-benzyloxyphenyl)-2-methoxyacrylic acid Na salt. The latter was hydrogenated in MeOH over [Ru(p-cymene)[(S)-dm-segphos]]Cl in MeOH at 5 MPa and 60° for 16 h to give Na 3-(4-hydroxyphenyl)-2-methoxypropionate in 20% yield and 92.9% enantiomeric excess.

ACCESSION NUMBER: 2005:490344 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 143:43684

TITLE: Process for preparation of optically active

3-(4-hydroxyphenyl) propionic acids by reaction of protected 4-hydroxybenzaldehydes and glycolic acid derivatives to give cinnamates and asymmetric

hydrogenation of the latter.

INVENTOR(S): Yokozawa, Tohru; Shimizu, Hideo; Fujiwara, Takahiro;

Ino, Yasunori

PATENT ASSIGNEE(S): Takasago International Corporation, Japan

SOURCE: PCT Int. Appl., 95 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PA	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION	ΝΟ.		D.	ATE	
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		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
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	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MΖ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
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		SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,
		ΝE,	SN,	TD,	TG												
EF	1687	250			A1		2006	0809		EP 2	004-	8194	90		2	0041	126
	R:	CH,	DE,	ES,	FR,	GB,	LI,	IE									
JF	2007	5122	22		T		2007	0517		JP 2	006-	5204	29		2	0041	126
US	US 20070142472				A1		2007	0621		US 2	006-	5787	44		2	0060	510
PRIORIT	US 20070142472 IORITY APPLN. INFO.:									JP 2	003-	3982	01	i	A 2	0031	127

OTHER SOURCE(S): CASREACT 143:43684; MARPAT 143:43684

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s 144 and 'chiral? cataly?'

129174 'CHIRAL'

19 'CHIRALS'

129179 'CHIRAL'

('CHIRAL' OR 'CHIRALS')

2 'CATALY'

0 'CHIRAL? CATALY?'

('CHIRAL'(W)'CATALY')

L47 0 L44 AND 'CHIRAL? CATALY?'

=> 147 and chiral catalyst

L47 IS NOT A RECOGNIZED COMMAND

The previous command name entered was not recognized by the system. For a list of commands available to you in the current file, enter "HELP COMMANDS" at an arrow prompt (=>).

=> s 144 and chiral catalyst

129174 CHIRAL

19 CHIRALS

129179 CHIRAL

(CHIRAL OR CHIRALS)

827404 CATALYST

823540 CATALYSTS

1060120 CATALYST

(CATALYST OR CATALYSTS)

2138 CHIRAL CATALYST

(CHIRAL (W) CATALYST)

L48 0 L44 AND CHIRAL CATALYST

=> s 144 and chiral

129174 CHIRAL

19 CHIRALS

129179 CHIRAL

(CHIRAL OR CHIRALS)

L49 24 L44 AND CHIRAL

=> s 124 and catalyst

12 "TOHRU"

5 "YOKOZAWA"

0 "TOHRU YOKOZAWA"

("TOHRU"(W)"YOKOZAWA")

827404 CATALYST

823540 CATALYSTS

1060120 CATALYST

(CATALYST OR CATALYSTS)

L50 0 L24 AND CATALYST

=> s 144 and chiral ligand

129174 CHIRAL

19 CHIRALS

129179 CHIRAL

(CHIRAL OR CHIRALS)

357859 LIGAND

243937 LIGANDS 486682 LIGAND

(LIGAND OR LIGANDS)

4290 CHIRAL LIGAND

(CHIRAL(W)LIGAND)

L56 2 L44 AND CHIRAL LIGAND

=> d 144 ibib abs 1-2

L44 ANSWER 1 OF 617 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2008:1522131 CAPLUS Full-text

TITLE: Inhibitory activity of Melissa officinalis L. extract

on Herpes simplex virus type 2 replication

AUTHOR(S): Mazzanti, G.; Battinelli, L.; Pompeo, C.; Serrilli, A.

M.; Rossi, R.; Sauzullo, I.; Mengoni, F.; Vullo, V.

CORPORATE SOURCE: Department of Human Physiology and Pharmacology,

"Sapienza" University, Rome, Italy

SOURCE: Natural Product Research, Part B: Bioactive Natural

Products (2008), 22(16), 1433-1440

CODEN: NPRPEA

PUBLISHER: Taylor & Francis Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

Melissa officinalis (Lamiaceae) (lemon balm) is used in folk medicine for AΒ nervous complaints, lower abdominal disorders and, more recently, for treating Herpes simplex lesions. In this work the antiviral activity of a hydroalcoholic extract of lemon balm leaves against the Herpes simplex virus type 2 (HSV-2) was assessed by the cytopathic effect inhibition assay on Vero cells (ATCC CCL-81), in comparison with acyclovir. The cytotoxicity of the extract on Vero cells was previously tested by evaluating the cellular death and was confirmed by the Trypan blue test. Lemon balm showed to reduce the cytopathic effect of HSV-2 on Vero cells, in the range of non-toxic concns. of 0.025-1 mg mL-1 (with reference to the starting crude herbal material). The maximum inhibiting effect (60%) was obtained with 0.5~mg mL-1. The viral binding assay showed that the extract does not prevent the entry of HSV-2 in the cells, thus suggesting a mechanism of action subsequent to the penetration of the virus in the cell. The extract was also chemical characterized by NMR and HPLC anal.; it showed to contain cinnamic acid-like compds., mainly rosmarinic acid (4.1% weight/weight). Our expts. support the use of lemon balm for treating Herpes simplex lesions and encourage clin. trials on this medicinal plant.

L44 ANSWER 2 OF 617 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2008:1433048 CAPLUS Full-text

DOCUMENT NUMBER: 150:41243

TITLE: Method for preparing salvianolic acid A of Salvia

miltiorrhiza with salvianolic acid B

INVENTOR(S): Li, Zhigang; Gu, Qun; Qu, Shoufeng; Mi, Changjiang;

Lin, Zhirong

PATENT ASSIGNEE(S): Beijing Bencao Tianyuan Pharmaceutical Research

Institute, Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 20pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

CN 101311160 A 20081126 CN 2007-10099618 20070525 PRIORITY APPLN. INFO.: CN 2007-10099618 20070525

The invention relates to a method for preparing salvianolic acid A of Salvia miltiorrhiza with salvianolic acid B. The method comprises (1) dissolving salvianolic acid B in water, adjusting pH to 3.5-6, heating at 110-130°C for 1-6h, filtering, concentrating, and drying to obtain salvianolic acid A extract; or dissolving salvianolic acid B in water, adjusting pH to 3.5-6, extracting under irradiation of microwave at 915-2450 MHZ and 1000-15000 W for 0.5-2 h, filtering, concentrating, and drying to obtain salvianolic acid A extract; (2) purifying by nonpolar or weak-polar macroporous resin column chromatog., silica gel chromatog., dextran gel LH-20 chromatog., polyamide chromatog. and/or extraction

L60 STRUCTURE UPLOADED

=> s 160

SAMPLE SEARCH INITIATED 10:37:49 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 597 TO ITERATE

100.0% PROCESSED 597 ITERATIONS 26 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 10475 TO 13405 PROJECTED ANSWERS: 215 TO 825

L61 26 SEA SSS SAM L60

=> d scan

> s 161/prep

122 L61

4705416 PREP/RL

L62 50 L61/PREP

(L61 (L) PREP/RL)

=>

=> s 165/prep

3709 L65

4705416 PREP/RL

L66 1056 L65/PREP

(L65 (L) PREP/RL)

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L67 600 L66 AND (PY<2003 OR AY<2003 OR PRY<2003)

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1145195 OPTICAL? 4457524 ACTIV?

51440 OPTICAL? ACTIV?

(OPTICAL?(W)ACTIV?)

L68 28 L67 AND (OPTICAL? ACTIV?)

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L69 52 L67 AND CATALY?

=> d 169 ibib abs 1-5

L69 ANSWER 1 OF 52 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:491168 CAPLUS Full-text

DOCUMENT NUMBER: 139:69049

TITLE: Preparation of substituted phenylpropionic acid

derivatives as agonists to human peroxisome proliferator-activated receptor alpha (PPAR)

INVENTOR(S): Lindstedt Alstermark, Eva-Lotte; Olsson, Anna

Christina; Li, Lanna

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PAT	CENT	NO.			KIN	D -	DATE				ICAT				D.	ATE		
WO	2003	0518	21		A1		2003	0626							2	0021	218	<
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							MD,											
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	2002						2004											
	2004						2005											
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	1620		1 1		A		2005			CN 2						0021		
_	2005						2005			JP 2	003-	5527	09		2	0021	218	<
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	5332						2006 2006	-		ıw ∠ NZ 2		-				0021	_	
	2558	-					2006 2006			NZZ						0021		
	3387				Б		2006			IW Z						0021		
	1896	_			A		2008			CN 2						0021	_	
CIA	1000	040			Λ		2001	O T T /		C14 Z	000-	1000	1113		۷	0021	0	` '

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ES 2271381
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          AT 363466
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MX 2004PA06004
NO 2004003023
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A 20060222 ZA 2004-4658

A 20040927 MX 2004-PA6004

A 20040715 NO 2004-3023
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                                                A1 20051222 US 2004-26806

A1 20070302 HK 2005-100831

A1 20050804 US 2005-499261

A 20051208 JP 2005-235794

A 20061102 JP 2006-123399
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          US 20050171204
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      JP 2003-552710
      A3 20021218 <--</td>

      WO 2002-GB5738
      W 20021218 <--</td>

      WO 2002-GB5744
      A 20021218 <--</td>

      GB 2002-29931
      A 20021221 <--</td>

      GB 2003-14079
      A 20030618

      WO 2003-GB5602
      A 20031219

      WO 2004-EP6597
      A 20040617

      US 2005-499261
      A2 20050304

PRIORITY APPLN. INFO.:
```

OTHER SOURCE(S): MARPAT 139:69049

GΙ

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/ Structure 43 in file .gra /
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AΒ The S enantiomer of I, n = 1 or 2, (C6H13 = hexyl) as well as their pharmaceutically acceptable salts, solvates, crystalline forms and prodrugs are synthesized using various solvents and in presence of charcoal-supported palladium catalyst. The utility of these compds. in clin. conditions such as lipid disorders (dyslipidemias) whether or not associated with insulin resistance and therapeutic and other pharmaceutical activities is also investigated. For example, (2S)-3-(4{2-[benzyl(hexyl)amino]-2oxoethoxy}phenyl)2-ethoxypropionic acid was prepared in 58% yield via reaction of (2S)-2-ethoxy-3-(4-hydroxyphenyl)propanoate and benzyl bromoacetate. THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 8 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L69 ANSWER 2 OF 52 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:22888 CAPLUS Full-text

DOCUMENT NUMBER: 138:73085

TITLE: Preparation of 3-aryl- α -oxy substituted

propanoic acids for treatment of type II diabetes and

related disorders

Potlapally, Rajender Kumar; Velagala, Venkata Rama INVENTOR(S):

Murali Krishna Reddy; Mamillapalli, Ramabhadra Sarma;

Gaddam, Om Reddy

Reddy's Research Foundation, India PATENT ASSIGNEE(S):

SOURCE: PCT Int. Appl., 54 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Pat.ent. LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

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PATENT NO.
                     KIND DATE APPLICATION NO.
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                                       _____
                      A1 20030109 WO 2001-IN124
    WO 2003002575
                                                            20010628 <--
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           GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
           LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
           RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US,
           UZ, VN, YU, ZA, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
           DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
           BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    AU 2001277663 A1 20030303 AU 2001-277663
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WO 2001-IN124 W 20010628 <--
                             20041209
PRIORITY APPLN. INFO.:
OTHER SOURCE(S): CASREACT 138:73085; MARPAT 138:73085
GΙ
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/ Structure 44 in file .gra /

AB 3-Aryl- α -oxy substituted propanoic acids [I; wherein R1 = tertbutyldimethylsilyl, trimethylsilyl, alkoxyalkyl; R2 = H, (substituted) (C1-C6) alkyl] were prepared For example, Me tert-butyldimethylsilyloxy-3-(4-hydroxyphenyl)propanoate (II) was prepared in three steps. The prepared compds. are useful for treating diabetes, obesity, glucose intolerance, insulin resistance and other related disorders such as hypertension, coronary heart disease, atherosclerosis, stroke, peripheral vascular diseases and related disorders (no data). The prepared compds. are also useful for reducing total cholesterol, body weight, blood plasma glucose, triglycerides, LDL, VLDL and free fatty acids (no data).

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L69 ANSWER 3 OF 52 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:927394 CAPLUS Full-text

DOCUMENT NUMBER: 138:4416

TITLE: Process for the preparation 3-aryl-2-hydroxypropionic

acid derivatives

INVENTOR(S): Ehrl, Robert; Ioannidis, Panagiotis; Mackintosh,

William

PATENT ASSIGNEE(S): Astrazeneca Ab, Swed. SOURCE: PCT Int. Appl., 21 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT	NO.			KIN	D :	DATE			APPL	ICAT	ION I	NO.		D	ATE	
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WO 2002	0968	65		A1		2002	1205	1	WO 2	002-	SE10	40		20	0020	530 <
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	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,
	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	NO,	NZ,	OM,	PH,
	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TN,	TR,	TT,	TZ,

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UA, UG, US, UZ, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     CA 2448658
                               20021205 CA 2002-2448658
                                                                  20020530 <--
                         Α1
     AU 2002309400
                         Α1
                                20021209
                                           AU 2002-309400
                                                                  20020530 <--
     NZ 529815
                         Α
                               20031219
                                           NZ 2002-529815
                                                                  20020530 <--
     EP 1404651
                         A1
                               20040407
                                           EP 2002-736372
                                                                  20020530 <--
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     BR 2002010125
                               20040608
                                          BR 2002-10125
                                                                  20020530 <--
                         Α
                         Τ
     JP 2004528388
                               20040916
                                           JP 2003-500045
                                                                  20020530 <--
     CN 1535262
                         Α
                               20041006
                                           CN 2002-814965
                                                                  20020530 <--
     CN 1247537
                         С
                               20060329
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                               20040916 ZA 2003-9216
                                                                  20031126 <--
                        Α
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                               20040227
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                                                                  20031128 <--
                        A1
                                           US 2004-479159
     US 20050014955
                               20050120
                                                                  20040823 <--
                                                               A 20010601 <--
PRIORITY APPLN. INFO.:
                                           SE 2001-1979
                                                               A 20020402 <--
                                           SE 2002-1004
                                                               W 20020530 <--
                                           WO 2002-SE1040
OTHER SOURCE(S):
                        CASREACT 138:4416; MARPAT 138:4416
     2-Ethoxy-3-[4-[2-(4-methanesulfonyloxyphenyl)ethoxy]phenyl]propanoic acid (I)
     or its alkyl esters were prepared by etherification of 2-ethoxy-3-(4-
     hydroxyphenyl)propanoic acid (II) or alkyl ester with 4-MeSO3C6H4CH2CH2-X (X
     is a suitable leaving group) in the presence of a base and a phase transfer
     catalyst at 50-150°C. Thus, 4-MeSO3C6H4CH2CH2O3SMe (1.01 mol), (S)-II Et
     ester (406 mmol) and PEG-400 (81 mmol) were melted together at 110 °C, Na2CO3
     (536 mmol) added under vigorous stirring, and the reaction continued at this
     temperature for 5.5 h. Saponification of the ester afforded (S)-I, a compound
     for therapeutic use in the Insulin Resistance Syndrome (IRS).
                              THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                         2
                              RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L69 ANSWER 4 OF 52 CAPLUS COPYRIGHT 2009 ACS on STN
                        2002:793403 CAPLUS Full-text
ACCESSION NUMBER:
DOCUMENT NUMBER:
                        137:310931
TITLE:
                        Preparation of phenylalkanoic acid derivatives as
                        preventive or remedial agents for digestive tract
                        diseases
                        Horizoe, Tatsuo; Shinoda, Masanobu; Emori, Eita;
INVENTOR(S):
                        Matsuura, Fumiyoshi; Kaneko, Toshihiko; Ohi, Norihito;
                        Kasai, Shunji; Yoshitomi, Hideki; Yamazaki, Kazuto;
                        Miyashita, Sadakazu; Hihara, Taro; Seiki, Takashi;
                        Clark, Richard; Harada, Hitoshi
PATENT ASSIGNEE(S):
                        Eisai Co., Ltd., Japan
```

PATENT	NO.			KIN	D :	DATE			APPL	ICAT	ION I	NO.		D	ATE	
WO 2002	W: AE, AG,					2002	1017	1	wo 2	002-	JP30	06		2	00203	327 <
W:	ΑE,	ΑG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,
	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TN,	TR,	TT,	TZ,

PCT Int. Appl., 344 pp.

CODEN: PIXXD2

Pat.ent.

Japanese

SOURCE:

LANGUAGE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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UA, UG, US, UZ, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
            CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
    AU 2002242989
                               20021021
                                           AU 2002-242989
                                                                  20020327 <--
                         Α1
                                                               A 20010330 <--
PRIORITY APPLN. INFO.:
                                           JP 2001-101465
                                           JP 2001-105131
                                                              A 20010403 <--
                                           WO 2002-JP3006
                                                              W 20020327 <--
```

OTHER SOURCE(S): MARPAT 137:310931

GΙ

/ Structure 45 in file .gra /

AΒ Disclosed is a preventive/remedy for digestive tract or inflammatory diseases, which contains as the active ingredient a novel carboxylic acid derivative represented by the following formula [I; R1 = H, OH, each (un)substituted C1-6 alkyl, C1-6 alkoxy, C1-6 alkylthio, C1-6 hydroxyalkyl, C1-6 hydroxyalkoxy, C1-6 hydroxyalkylthio, C1-6 aminoalkyl, C1-6 aminoalkoxy, C1-6 aminoalkylthio, C2-12 alkoxyalkyl, C3-7 cycloalkyl, C3-7 cycloalkyloxy, C3-7 cycloalkylthio, C2-6 alkenyl, C2-6 alkenyloxy, or C2-6 alkenylthio, etc.; L = a single or double bond, each (un)substituted C1-6 alkylene, C2-6 alkenylene, or C2-6 alkynylene; M = a single bond, each (un)substituted C1-6 alkylene, C2-6 alkenylene, or C2-6 alkynylene; T = a single bond, each (un)substituted C1-3alkylene, C2-3 alkenylene, or C2-3 alkynylene; W = 2,4-dioxothiazolidin-5-yl,2,4-dioxothiazolidin-5-ylidene, carboxy, (un)substituted CONH2; X = O, (un) substituted C2-6 alkenylene, hydroxymethylene, CO, CS, N-(un) substituted CQNH, NHCQ, SO2NH, NHSO2, or NHCQNH (Q = O, S); Y = (un) substituted C5-12 aromatic hydrocarbyl or C3-7 aliphatic hydrocarbyl optionally containing ≥1 heteroatoms; ring Z = C5-6 aromatic hydrocarbyl; Y = (un) substituted aromatic hydrocarbon group optionally containing ≥1 heteroatoms; some provisos given], a salt of the derivative, or a hydrate of either. The above digestive tract diseases include (1) inflammatory digestive tract diseases such as ulcerous colitis, Crohn's disease, pancreatitis, and gastritis, (2) digestive tract proliferative diseases such as digestive tract benign rumors, digestive tract polyp, hereditary (genetic) polyposis syndromes, colon cancer, rectum cancer, and stomach cancer, and (3) digestive tract ulcerous diseases such as duodenal ulcer, stomach ulcer, esophagus ulcer, regurgitant esophagitis, stress ulcer or erosion, erosion caused by drugs, and Zollinger-Ellison syndromes. The above inflammatory diseases include arthritic rheumatism, multiple sclerosis, immunodeficiency, cachexia, osteoarthritis, osteoporosis, asthma, and allergy. The compds. I are triple agonists for PPAR (peroxisome proliferator-activated receptor) α , β , and γ subtype. Thus, 2-isopropoxy-3-[4-methoxy-3-[[4-(trifluoromethyl)benzyl]amino]carbonyl]phenyl]propanoic acid in vitro showed the transcription activity for PPARlpha, eta, and γ with EC50 of 0.08, 2.513, and 0.382 μ M, resp., in CV-1 cell. (2S)-3-[3-[[(2,4-dichlorobenzoyl)amino]methyl]-4-methoxyphenyl]-2- isopropoxypropanoic acid at 1 mg/kg/day p.o. for 3 days showed a disease activity index based on diarrhea, bloody excrement, and weight loss (DAI) of 2.0±0.3 in mice suffering from colitis induced by dextran sulfate sodium salt vs. 2.8 ± 0.2 for the control group and 2.1 ± 0.3 for the mice treated with rosiglitazone at 30 mg/kg/day. Many compds. prepared do not possess the thiazolidine skeleton and thereby may completely avoid toxicity such as liver disorder which was noted in the past as a problem for compds. having PPARy agonist activity.

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
ACCESSION NUMBER:
                         2002:620035 CAPLUS Full-text
DOCUMENT NUMBER:
                         138:89669
TITLE:
                         Efficient synthesis of antihyperglycemic
                          (S) -\alpha-aryloxy-\beta-phenylpropionic acid
                         derivative using a bifunctional asymmetric
                         catalvst
AUTHOR(S):
                         Takamura, Makoto; Yanagisawa, Hiroaki; Kanai, Motomu;
                         Shibasaki, Masakatsu
CORPORATE SOURCE:
                         Medicinal Chemistry Research Laboratories, Sankyo Co.,
                         Ltd., Tokyo, 140-8710, Japan
SOURCE:
                         Chemical & Pharmaceutical Bulletin (2002),
                         50(8), 1118-1121
                         CODEN: CPBTAL; ISSN: 0009-2363
PUBLISHER:
                         Pharmaceutical Society of Japan
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
OTHER SOURCE(S):
                         CASREACT 138:89669
GΙ
L70
        STRUCTURE UPLOADED
=> s sss full 170
FULL SEARCH INITIATED 11:17:13 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 12087 TO ITERATE
100.0% PROCESSED
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L71
           7767 SEA SSS FUL L70
=> s 171/prep
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L72
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       4502889 AY<2003
       3971612 PRY<2003
           600 L72 AND (PY<2003 OR AY<2003 OR PRY<2003)
L73
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       1500302 CATALY?
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L74
=> s 173 and ('asymmetric hydrogenation' or ' ?select? hydrogenation')
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('ASYMMETRIC' OR 'ASYM')

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             0 L73 AND ('ASYMMETRIC HYDROGENATION' OR ' ?SELECT? HYDROGENATION'
=> d 174 ibib abs 1-4
L74 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2001:836609 CAPLUS Full-text
DOCUMENT NUMBER:
                         136:118291
                         Total synthesis of biologically active compounds
TITLE:
                         related to plant disease and the physiological
                         function
                         Toshima, Hiroaki
AUTHOR(S):
                        Department of Bioresource Science, College of
CORPORATE SOURCE:
                         Agriculture, Ibaraki University, Ibaraki, 300-0393,
SOURCE:
                         Yuki Gosei Kagaku Kyokaishi (2001), 59(11),
                         1121-1129
                         CODEN: YGKKAE; ISSN: 0037-9980
PUBLISHER:
                         Yuki Gosei Kagaku Kyokai
                         Journal; General Review
DOCUMENT TYPE:
LANGUAGE:
                         English
     A review, total synthesis of biol. active compds. related to plant disease and
     the physiol. function has been accomplished. Coronatine, its related compds.,
     \beta-resorcylic acid derivs., decumbic acid, aliphatic \delta-lactones, cepaciamides,
     and piscidic acids, were selected as the synthetic targets. In these
     syntheses, the chiral pool method and catalytic asym. synthesis were also
     applied to introduce the requisite stereogenic centers. Combination of the
     two methods made it possible to synthesize a sufficient amount of the required
     enantiomers and diastereomers for biol. studies.
REFERENCE COUNT:
                         43
                               THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L74 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2009 ACS on STN
                         1999:784755 CAPLUS Full-text
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         132:166053
                         Total syntheses of all four stereoisomers of piscidic
TITLE:
                         acid via catalytic asymmetric
                         dihydroxylation of (Z)- and (E)-trisubstituted olefins
                         Toshima, Hiroaki; Saito, Masatoshi; Yoshihara,
AUTHOR(S):
                         Teruhiko
                         Department of Bioscience and Chemistry, Faculty of
CORPORATE SOURCE:
                         Agriculture, Hokkaido University, Sapporo, 060-8589,
                         Bioscience, Biotechnology, and Biochemistry (
SOURCE:
                         1999), 63(11), 1934-1941
```

CODEN: BBBIEJ; ISSN: 0916-8451

PUBLISHER: Japan Society for Bioscience, Biotechnology, and

Agrochemistry

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 132:166053

Four stereoisomers of (2S,3R)-(+)-piscidic acid were synthesized with high optical purity via Sharpless catalytic asym. dihydroxylation of (Z)- and (E)-trisubstituted olefins in 6 steps from (4-hydroxyphenyl)pyruvic acid. That is Wittig reaction of Me (4-hydroxyphenyl)pyruvate with (carbomethoxymethylene)triphenylphosphorane give (Z)- and (E)-trisubstituted olefins in a 3:1 ratio, protecting the phenolic hydroxyl group with tert-butyldimethylsilyl, then the (Z)-olefin was subjected to asym. dihydroxylation by using the chiral ligand, dihydroquinidine 1,4-anthraquinonediyl diether, gave the product with 89% e.e. Desilylation and subsequent alkaline hydrolysis gave (2S,3R)-(+)-piscidic acid with > 99% e.e. after recrystn. Use of ligand, dihydroquinine 1,4-anthraquinonediyl diether, gave (2R,3S)-(-)-piscidic acid. Asym. dihydroxylation of the (E)-olefin with phthalazine ligands (dihydroquinidine and dihydroquinine 1,4-phthalazinediyl diethers) also gave high e.e. values product, followed by the same procedure mentioned

above, gave (2S,3S)-(+)-3-epi-piscidic acid and (2R,3R)-(-)-2-epi-piscidic

acid resp.

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L74 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1999:363435 CAPLUS Full-text

DOCUMENT NUMBER: 131:184794

TITLE: Total synthesis of (+)-(2S,3R)-Piscidic acid via

catalytic asymmetric dihydroxylation of a

trisubstituted olefin

AUTHOR(S): Toshima, Hiroaki; Saito, Masatoshi; Yoshihara,

Teruhiko

CORPORATE SOURCE: Department of Bioscience and Chemistry, Faculty of

Agriculture, Hokkaido University, Sapporo, 060-8589,

Japan

SOURCE: Bioscience, Biotechnology, and Biochemistry (

1999), 63(5), 964-967

CODEN: BBBIEJ; ISSN: 0916-8451

PUBLISHER: Japan Society for Bioscience, Biotechnology, and

Agrochemistry

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 131:184794

AB (+)-(2S,3R)-Piscidic acid was efficiently synthesized with high optical purity (90% e.e.) via Sharpless catalytic asym. dihydroxylation of a trisubstituted olefin in only 6 steps from com. available 4-hydroxyphenyl-pyruvic acid as the starting material. The reaction proceeded with high optical purity by using the chiral ligands, dihydroquinidine 2,5-diphenyl-4,6-pyrimidinediyl diether or dihydroquinidine 1,4-anthraquinonediyl diether.

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L74 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1992:426523 CAPLUS Full-text

DOCUMENT NUMBER: 117:26523

ORIGINAL REFERENCE NO.: 117:4791a,4794a

TITLE: Enantioselective synthesis of calcium channel blockers

of the diltiazem group

AUTHOR(S): Schwartz, Alan; Madan, Pradeep B.; Mohacsi, Erno;

O'Brien, Jay P.; Todaro, Louis J.; Coffen, David L. CORPORATE SOURCE: Roche Res. Cent., Hoffmann-La Roche Inc., Nutley, NJ,

07110, USA

SOURCE: Journal of Organic Chemistry (1992), 57(3),

851 - 6

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 117:26523

GΙ

/ Structure 47 in file .gra /

AB A lipase-catalyzed kinetic resolution of racemic trans-2-phenylcyclohexanol readily provides the (-)-(1R,2S) enantiomer. This alc. is employed as its chloroacetate in a chiral auxiliary-induced asym. Darzens glycidic ester condensation with p-anisaldehyde. Crystallization of the Darzens product affords enantiomerically pure phenylcyclohexyl (methoxyphenyl)glycidate I, the structure of which was established by x-ray crystallog. The use of this glycidic ester in syntheses of diltiazem II (R = R1 = H) and maltiazem II [RR1 = (CH:CH)2], members of the diltiazem group of calcium channel blockers, provides these drug substances directly in enantiomerically pure form.

=> d 173 ibib abs 1-5

L73 ANSWER 1 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2008:994060 CAPLUS Full-text

DOCUMENT NUMBER: 149:306485

TITLE: Di-m-Chlorobis[Bis-(cyclooctene)rhodium]

AUTHOR(S): Judd, Andrew S.

CORPORATE SOURCE: USA

SOURCE: e-EROS Encyclopedia of Reagents for Organic Synthesis

(2001), No pp. given. John Wiley & Sons,

Ltd.: Chichester, UK.

CODEN: 69KUHI

URL: http://www3.interscience.wiley.com/cgi-

bin/mrwhome/104554785/HOME

DOCUMENT TYPE: Conference; General Review; (online computer file)

LANGUAGE: English

OTHER SOURCE(S): CASREACT 149:306485

AB A review of the article Di-m-Chlorobis[Bis-(cyclooctene)rhodium].

L73 ANSWER 2 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2008:302246 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 148:262617

TITLE: Preparation of pyrimidine- and triazine-derivative

endothelin receptor antagonists

INVENTOR(S): Riechers, Hartmut; Klinge, Dagmar; Amberg, Wilhelm;

Kling, Andreas; Mueller, Stefan; Baumann, Ernst; Rheinheimer, Joachim; Vogelbacher, Uwe Josef; Wernet,

Wolfgang; Unger, Liliane; Raschack, Manfred

PATENT ASSIGNEE(S): Abbott Gmbh & Co. KG, Germany

SOURCE: U.S., 18pp., Cont. of U.S. Ser. No. 748,184.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

	ATENT NC								AE 									
U	S 710920 S 200400	5			B2	_		0919								 20030		
U	S 200400	92	742		A1		2004	0513										
D	E 195330	23			A1		1996	0418	DE	1	995-	1953	3023			19950	907	<
	E 195330							0516										
W	0 961191	4			A1		1996	0425	WC) 1	995-	EP39	63			19951	007	<
	W: A	U,	BG,	BR,	BY,	CA	, CN,	CZ,	FI, F	ΙU,	JP,	KR,	KΖ,	MX,	ИС	, NZ,	PL,	
		,	,	,	SK,													
									GB, G									
	P 111095								EF	2	001-	1038	89			19951	007	<
E	P 111095							0929										
	R: A																	
	S 593273								US									
	S 596913				Α				US									
	S 619795				В1				US									
	S 200200							0502		2	000-	7481	84			20001	227	<
	S 660004				В2			0729										
	S 200601							0720		3 2	006-	3778	79			20060	316	<
	S 711909				В2			1010										
	S 200602							1207	US	3 2	006-	5022	57			20060	810	<
	S 200602							1207								20060		
_	S 200702				A1		2007	0830								20070		
PRIORI	TY APPLN	•	INFO	.:									851			19941	_	
													3023			19950	-	
													63			19951		
													99			19970		
													52		_	19981	_	
													70			19990		
													84			20001		
													16		-	19951		<
													75			20030		
OTHER	SOURCE (S	٧.			MAN	- A - F	1.40	2020		5 2	006-	5022	57		RT	20060	8T0	
UTHER	SUITELETS	. 1 *			MARI	- 4	148 *	/h/h	1 /									

OTHER SOURCE(S): MARPAT 148:262617

GI

/ Structure 48 in file .gra /

The title compds. I [R = CHO, tetrazolyl, CN, CO2H, groups cleavable to CO2H; R2 = (un)substituted NH2, halogen, (un)substituted alkyl, etc.; R3 = H, OH, (un)substituted NH2, halogen, (un)substituted alkyl, etc.; R4, R5 = (un)substituted Ph or naphthyl; R6 = H, alkyl, alkenyl, alkynyl, alkylcarbonyl, (un)substituted Ph, etc.; X = N, (un)substituted CH; Y = direct bond, S, O; Z = S, O, SO, SO2, direct bond], and their pharmaceutically acceptable salts, are prepared and disclosed as endothelin receptor antagonists. In receptor binding assays, pyrimidine derivative II (R2 and R3 = MeO), m.p. 167°, demonstrated a Ki ETA of 6 nM. In particular, the racemate and individual enantiomers of II (R2 and R3 = Me) are claimed.

REFERENCE COUNT: 45 THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER: 2007:899716 CAPLUS Full-text

DOCUMENT NUMBER: 148:495971

TITLE: Process for the preparation of antidiabetic compound

PATENT ASSIGNEE(S): Dr. Reddy's Laboratories Ltd., India

SOURCE: Indian Pat. Appl., 30pp.

CODEN: INXXBQ

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

IN 2002MA00399 A 20070803 IN 2002-MA399 20020523 <-
PRIORITY APPLN. INFO.: IN 2002-MA399 20020523 <--

OTHER SOURCE(S): CASREACT 148:495971; MARPAT 148:495971

GΙ

/ Structure 49 in file .gra /

AB A invention relates to a process for the preparation of tromethamine salt of (-)-3-[4-(2-(5-ethyl-1-methyl-7-oxo-3-propyl-6,7-dihydro-1H-pyrazolo[4,3-d]pyrimidin-6-yl)ethoxy)phenyl]-2-alkoxypropionic acid of the formula I, where R is C1-6 alkyl, c. Example compound I (R = Et) was prepared by etherification of iso-Pr 3-(4-hydroxyphenyl)-2(S)-hydroxypropionate with 5-ethyl-6-(2-chloroethyl)-1-methyl-3-propyl-6,7-dihydro-1H-pyrazolo[4,3-d]pyrimidin-7-one; the resulting (-)-[((pyrazolo[4,3-d]pyrimidin-7-yl)ethoxy)phenyl]-2-hydroxypropanoate derivative underwent O-alkylation followed by hydrolysis to give the corresponding 2-alkoxypropanoic acid, which was reacted with tromethamine to give compound I.

L73 ANSWER 4 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2007:544712 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 148:449648

TITLE: An improved process for the preparation of phenoxazine

antidiabetic compounds

INVENTOR(S): Rao, Siripragada Mahender; Reddy, Chepyala Naveen

Kumar; Reddy, Challa Maheedhara; Sarma, Mamillapalli

Ramabhandra; Reddy, Gaddam Om

PATENT ASSIGNEE(S): Reddy's Laboratories Ltd., India

SOURCE: Indian Pat. Appl., 18pp.

CODEN: INXXBQ

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

IN 2002MA00508 A 20070511 IN 2002-MA508 20020708 <-
PRIORITY APPLN. INFO.: IN 2002-MA508 20020708 <--

OTHER SOURCE(S): CASREACT 148:449648; MARPAT 148:449648

GI

The invention relates to an improved process for the preparation of antidiabetic phenoxazinylethoxyphenylalkoxypropanoate arginine salts I [R1 = Me, Et, Pr, i-Pr]. The process uses inexpensive chems. and an easy resolution via chiral amine salts. Thus, condensation of 4-[2-(phenoxazin-10-yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the corresponding glycidic ester, which was hydrogenated over Pd/C, O-ethylated with di-Et sulfate in xylene, and saponified with aqueous NaOH in MeOH to give acid (±)-II. Resolution of this racemate using (-)-ephedrine and salification with L-arginine in i-PrOH gave I (R1 = Et).

L73 ANSWER 5 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2007:544708 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 148:449647

TITLE: Process for the preparation of antidiabetic

(S)-3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2-

ethoxypropanoic acid L-arginine salt

INVENTOR(S): Reddy, Gaddam Om

PATENT ASSIGNEE(S): Reddy's Laboratories Ltd., India

SOURCE: Indian Pat. Appl., 9pp.

CODEN: INXXBQ

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND APPLICATION NO. DATE DATE ______ _____ ____ _____ _____ IN 2002MA00017 20070511 IN 2002-MA17 20020107 <--PRIORITY APPLN. INFO.: IN 2002-MA17 20020107 <--CASREACT 148:449647; MARPAT 148:449647 OTHER SOURCE(S):

GI

/ Structure 51 in file .gra /

AB Claimed is a process for the preparation of compound I [R1 = alkyl] comprising (a) reacting a [(phenoxazinyl)ethoxy]phenylpropanoic acid derivative with 1-phenylethylamine in the presence of a solvent at 20°C to 50°C, (b) reacting the resulting salt with L-arginine in the presence of a solvent in the range of 20°C to reflux temperature for 4 to 24 h; (c) isolating the resulting product. I is an antidiabetic agent (no data). Thus, (S)-3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-2- ethoxypropanoic acid (II) in Et acetate was reacted with 1-phenylethylamine to give a salt; a solution of said salt in methanol was treated with L-arginine to give II L-arginine salt.

L73 ANSWER 6 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2007:183455 CAPLUS Fuil-text

DOCUMENT NUMBER: 146:462276

TITLE: Preparation of pyrimidinone derivatives for use in

medicine and pharmaceutical compositions containing

them

INVENTOR(S): Madhavan, Gurram Ranga; Venkateswarlu, Akella;

Rajagopalan, Ramanujam; Chakrabarti, Ranjan; Misra, Parimal; Lohray, Braj Bhushan; Lohray, Vidya Bhushan;

Rao, Paraselli Bheema

PATENT ASSIGNEE(S): Dr. Reddy's Research Foundation, India

SOURCE: Indian Pat. Appl., 112pp.

CODEN: INXXBQ

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

-----IN 2001MA00568 A 20050304 IN 2001-MA568 20010710 <-
PRIORITY APPLN. INFO: IN 2001-MA568 20010710 <--

OTHER SOURCE(S): CASREACT 146:462276; MARPAT 146:462276

GΙ

/ Structure 53 in file .gra /

AΒ Title compds. I [X = O or S; R1-3 independently when attached to C = H, halo,OH, CN, etc.; when R3 is attached to N, R3 = H, OH, CHO, etc.; R4 = H, halo, alkyl, etc.; R5 = H, halo, alkoxy, etc.; or R4 and R5 join to form a bond; R6 = H, (un)substituted alkyl, acyl, aryl, etc.; R7 = H, (un)substituted alkyl, cycloalkyl, aryl, etc.; Y = O or NR8, where R8 = H, alkyl, aryl, etc.; or R7 and R8 may join to form cyclic or heterocyclic ring; n = 1-4; Ar = (un) substituted aryl or heterocyclic group], and their pharmaceutically acceptable salts, are prepared and disclosed as antiobesity and anticholesteremic agents. Thus, e.g., II was prepared by substitution of Et 2-ethoxy-3-[4-(2-haloethoxy)phenyl]propanoate with 2-ethyl-4-phenyl-1,6dihydropyrimidin-6-one followed by hydrolysis to provide the carboxylic acid. I were evaluated for hPPARy activity, e.g., II was determined to have a PPARy value of 23.9 at 1 μ M. The present invention relates to novel antiobesity and hypocholesterolemic compds., their derivs., their analogs, their tautomeric forms, their stereoisomers, their polymorphs, their pharmaceutically acceptable salts, their pharmaceutically acceptable solvates and pharmaceutically acceptable composition containing them.

L73 ANSWER 7 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2007:181507 CAPLUS Full-text

DOCUMENT NUMBER: 148:355830

TITLE: Preparation of thiodiphenylamines and related

compounds as hypolipemic agents

INVENTOR(S): Lohray, Braj Bhushan; Lohray, Vidya Bhushan; Bajji,

Ashok Channa Veerappa; Kalchar, Shivaramayya; Rao, Paraselli Bheema; Madhavan, Gurram Ranga; Rajacopalan,

Ramanujam; Chakrabarti, Ranjan

PATENT ASSIGNEE(S): Dr. Reddy's Research Foundation, India

SOURCE: Indian Pat. Appl., 75pp.

CODEN: INXXBQ

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
IN 1999MA00483	А	20050304	IN 1999-MA483	19990428 <
PRIORITY APPLN. INFO.:			IN 1999-MA483	19990428 <

GI

/ Structure 54 in file .gra /

AB Title compds. I [Z = (CH2)m; m = 1-4; n = 0-1; R1, R2, R3, R4 = H, halo, OH, etc.; A = 5 or 6-membered cyclic structure with provisos; X = 0, S, NR9; R9 = H, alkyl, aryl, etc.; Ar = fused aromatic, heterocyclic, etc.; R5 = H, OH, halo, etc.; R6 = H, OH, halo, etc.; R7 = H, alkyl, cycloalkyl, etc.; R8 = H, alkyl, cycloalkyl, etc.; Y = 0, NR10; R10 = H, alkyl, aryl, etc.] and their pharmaceutically acceptable salts were prepared For example, condensation of aldehyde II and triethyl-2-ethoxyphosphonoacetate afforded thiodiphenylamine III. In plasma triglyceride assays, 3-examples of compds. I exhibited 6-58% lowering of triglyceride concentration at 1 mg/kg dosage.

L73 ANSWER 8 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2005:1335635 CAPLUS Full-text

DOCUMENT NUMBER: 144:69628

TITLE: Preparation of phenoxyacetamide derivatives as

modulators of peroxisome proliferator-activated

receptors (PPAR)

INVENTOR(S): Lindstedt Alstermark, Eva-Lotte; Olsson, Anna

Christina; Li, Lanna

PATENT ASSIGNEE(S): Swed.

SOURCE: U.S. Pat. Appl. Publ., 47 pp., Cont.-in-part of U.S.

Ser. No. 499,261.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

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                                                                 A3 20040617
OTHER SOURCE(S): CASREACT 144:69628; MARPAT 144:69628
GΙ
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AΒ The phenyl-, phenoxy-, or phenylthioalkanamidetitle compds., (in particular phenoxyacetamide derivs.) (I) [A is situated in the ortho, meta or para position and represents CR3R4CR1R2COR, CR3:CR1COR (wherein R = H, alkyl, (un) substituted HO or NH2; R1 = alkyl, aryl, alkenyl, alkynyl, or when A is CR3R4CR1R2COR, R1 can also be cyano, (un) substituted HO, SH, OCONH2, SO2NH2, CO2H, etc.; R2 = H, halogen, alkyl, aryl, alkylaryl; R3, R4 = H, alkyl, aryl, alkylaryl); Y = 0, S, a single bond; n = an integer of 1-4; X = alkyl; R5, R6 = H, each (un)substituted C1-13 alkyl, C2-10 alkenyl, or C2-10 alkynyl; or R5, R6 = each (un)substituted C3-8 cycloalkyl, C3-C8 cycloalkenyl, aryl, heterocyclyl, or heteroaryl; or R5 and R6 together with the nitrogen atom to which they are attached form a single or a fused heterocyclic system] are prepared These compds. are useful in treating clin. conditions including lipid disorders (dyslipidemias) whether or not associated with insulin resistance, and other manifestations of the metabolic syndrome. Thus, a solution of 0.598 g N-butyl-N-[2-fluoro-4-(trifluoromethyl)benzyl]amine and 0.593 q [4-((2S)-2,3-diethoxy-3-oxopropyl)phenoxy]acetic acid in 20 mL CH2Cl2 was treated with 0.80 mL N, N-diisopropylethylamine and 0.674 g O-(benzotriazol-1-yl)-N,N,N',N'-tetramethyluronium tetrafluoroborate and the reaction mixture was stirred at room temperature overnight to give, after workup and silica gel chromatog., 74% Et (2S)-3-[4-[2-[buty1[2-fluoro-4-(trifluoromethyl)benzyl]amino]-2- oxoethoxy]phenyl]-2-ethoxypropanoate (II). A solution of 0.748 g II in 70 mL MeCN was treated with 35 mL 0.10 M LiOH and the reaction mixture was stirred at room temperature overnight, neutralized with 5% HCl, concentrated, acidified with 5% HCl, and extracted with EtOAc to qive 97% (2S)-3-[4-[2-[buty1[2-fluoro-4-(trifluoromethy1)benzy1]amino]-2oxoethoxy]phenyl]-2-ethoxypropanoic acid (III). III showed EC50 of 0.001 μ mol/L for human PPAr α .

L73 ANSWER 9 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2005:519176 CAPLUS Full-text

DOCUMENT NUMBER: 143:71779

TITLE: Red-rooted Salvia root anti-peptic ulcer effective

component and preparing process thereof

INVENTOR(S): Li, Hequan; Wang, Yulin; Li, Xi
PATENT ASSIGNEE(S): China Medical Univ., Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, No pp.

given

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1508133	A	20040630	CN 2002-144771	20021213 <
CN 1208330	С	20050629		
PRIORITY APPLN. I	INFO.:		CN 2002-144771	20021213 <

The present invention relates to effective region of salvia root for resisting peptic ulcer and its preparation process. The chemical composition of said effective region contains danshenphenolic acid component using danshenphenolic acid B as main substance and small quantity of fatty acid component. Its preparation process includes the following steps: decocting Chinese medicinal material salvia root by adding water, making filtrate pass through polyamide column, washing with distilled water to remove impurity, eluting with Et alc., reduced pressure recovering Et alc. solution, heating with water and dissolving the obtained red tan solid, precipitation, filtering to obtain yellow supernatant fluid, precipitating, washing with water, reduced pressure removing water content to obtain the product (DSE-F), brown color, its yield

is about 0.6%. Said product can be made into tablet, powder, capsule and decoction preparation $\ \ \,$

L73 ANSWER 10 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2005:483308 CAPLUS Full-text

DOCUMENT NUMBER: 143:225593

TITLE: Method for preparing salvianolic acid B of Salvia

miltiorrhiza

INVENTOR(S): Zhang, Fengxia; Wang, Jiaping; Tan, Xuebing

PATENT ASSIGNEE(S): Nanjing Hongqiao Institute of Medical Technology,

Peop. Rep. China

SOURCE: Faming Zhuanli Shenging Gongkai Shuomingshu, 13 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PRIORITY APPLN. INFO.:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1425659	A	20030625	CN 2002-160771	20021231 <
CN 1164582	С	20040901		

The method comprises extracting with water, acidifying to pH 1-6, purifying on macroporous resin column (ion exchange resin, Mcigel-Chp-20P, Sephadex LH-20, C18 bonded stationary phase, or C8 bonded stationary phase), and drying. The extraction and acidification processes may be simultaneously carried out by extraction with acidic solution (pH 1-6). The solvent extraction may be substituted by percolation, ultrasonic wave-aided extraction, or microwave-

aided extraction methods.

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L73 ANSWER 11 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2005:483216 CAPLUS Full-text

DOCUMENT NUMBER: 143:13395

TITLE: Medical composition of active constituent of

traditional Chinese medicine for treating

cardiovascular and cerebrovascular diseases and its

CN 2002-160771

20021231 <--

preparation

INVENTOR(S): Zhang, Weidong; Su, Juan; Zhang, Chuan

PATENT ASSIGNEE(S): Shanghai Botai Medical Science and Technology Co.,

Ltd., Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 10 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

CN 1425430 Α 20030625 CN 2002-155004 20021219 <--CN 100346798

С 20071107

PRIORITY APPLN. INFO.: CN 2002-155004 20021219 <--

The metal composition is composed of 1 part total salvianolic acid (its salvianolic acid B Mg salt content >50%) and 0.1-1 part total saponin (its astragalin A content >20%) of Astragalus membranaceus. The total salvianolic acid is isolated from Salvia multiorrhiza by percolating in acetone and purifying on macroporous adsorbent resin column. The total saponin is isolated from A. membranaceus by extracting with water and then purifying on macroporous adsorbent resin column. The metal composition may be used to prepare injection, large-capacity injection, powder injection, tablet, capsule, powder, etc.

L73 ANSWER 12 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:1154649 CAPLUS Full-text

DOCUMENT NUMBER: 142:93514

TITLE: Preparation of phenylpropanoic acid derivatives as

PPARα agonists

INVENTOR(S): Li, Lanna; Lindstedt-Alstermark, Eva-Lotte; Olsson,

Christina

PATENT ASSIGNEE(S): Astrazeneca Ab, Swed. SOURCE: PCT Int. Appl., 100 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

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CN	1835	913			Α		2006	0920		CN 2	004-	8002	3304		2	0040	617	
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CN	1012	3992	8		A		2008	0813		CN 2	-800	1000	9615		2	0040	617	
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MX 2005PA13712	A	20060308	MX	2005-PA13712		20051215	
JP 200 6 29 8 925	A	20061102	JΡ	2006-139673		20060519	
US 2006025 8866	A1	20061116	US	2006-477168		20060628	
US 20070244198	A1	20071018	US	2007-772474		20070702	
IN 2008DN04240	A	20080815	IN	2008-DN4240		20080519	
PRIORITY APPLN. INFO.:			GB	2003-14079	A	20030618	
			SE	2001-4334	Α	20011219 <	
			WO	2002-GB5738	\mathbb{W}	20021218 <	
			WO	2002-GB5744	Α	20021218 <	
			GB	2002-29931	A	20021221 <	
			WO	2003-GB5602	A	20031219	
			CN	2004-80023304	A3	20040617	
			EP	2004-740044	A3	20040617	
			JΡ	2006-515989	A3	20040617	
			WO	2004-EP6597	\mathbb{W}	20040617	
			IN	2005-DN5470	A3	20041128	
			US	2005-518777	A3	20050303	
			US	2005-499261	A2	20050304	
			US	2006-477168	A 1	20060628	

OTHER SOURCE(S): MARPAT 142:93514

GI

/ Structure 56 in file .gra /

AB Title compds. represented by the formula I [wherein A = CR3(R4)CR1(R2)COR or C(R3):C(R1)COR; R = H, alkoxy, (alkyl)aryloxy, amino, etc.; R1 = alkyl, aryl, alkenyl, alkynyl, etc.; R2 = H, halo, alkyl, (alkyl)aryl; R3, R4 = independently H, alkyl, (alkyl)aryl; T = O, S or a single bond; n = 1-4; R5, R6 = independently selected substituent comprising C, H, N, O, S, Se, P or halo; with provisos; optical isomers and racemates thereof as well as pharmaceutically acceptable salts, prodrugs, solvates and crystalline forms thereof] were prepared as PPARα agonists. For example, II was given in a multi-step synthesis starting from the reaction of 2,4-difluorobenzylamine with octanoic acid. I had EC50 values of less than 0.1 μmil/L for PPARα and showed the ration of the EC50(PPARγ) with EC50(PPARα) is greater than 150:1. Thus, I and their pharmaceutical compns. are useful for the treatment of clin. conditions including lipid disorders (dyslipidemias) whether or not associated with insulin resistance (no data).

REFERENCE COUNT: 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 13 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:918600 CAPLUS Full-text

DOCUMENT NUMBER: 142:113816

TITLE: Rosmarinic acid derivatives as an immunosuppressant

and an inhibitor of SH2

INVENTOR(S): Kang, Mi Ae; Kim, Hong Tae; Ko, Jae Gyun; Lee, Geon

Hyeong; Lee, Jong Seong; Oh, Jae Taek; Oh, Jong Eun;

Park, Si Hyeong; Won, Jong Hwa; Yoon, Yeong Dae

PATENT ASSIGNEE(S): Mogam Biotechnology Institute, S. Korea

SOURCE: Repub. Korean Kongkae Taeho Kongbo, No pp. given

CODEN: KRXXA7

DOCUMENT TYPE: Patent LANGUAGE: Korean

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
KR 2002006643	A	20020124	KR 2000-39730	20000711 <
PRIORITY APPLN. INFO.:			KR 2000-39730	20000711 <

AΒ A rosemarine derivative useful as an immunosuppressant and an inhibitor of SH2(src homol. region 2) in T lymphocyte cell kinase is provided, thereby diagnosing and preventing transplant rejection, inflammatory diseases, and autoimmune related diseases. The rosemarine derivative useful as an immunosuppressant and an inhibitor of SH2(src homol. region 2) in T lymphocyte cell kinase is represented by formula(1), in which R1, R2, R3, R4 and R5 are independent, wherein at least one of them is hydroxy, and others are selected from hydrogen, halogen, C1 to C3 alkoxy, aldehyde, carboxyl, amino, trifluormethyl and nitro; R6, R7, R8, R9 and R10 are independent, wherein at least one of them is hydroxy, and others are selected from hydrogen, halogen, C1 to C3 alkoxy, aldehyde, carboxyl, amino, trifluormethyl and nitro; X1 is O, S, NH, N-CH3, N-CH2CH3 or NHNH; X2 is CH2; X3 is (CH2)m; Y1 is selected from hydrogen, CH2, hydrogen, linear or branched alkyl or aryl-substituted amine; Y2 is none or -NZ11Z12, -O-Z2 or -S-Z2, wherein Z11 or Z12 is individually hydrogen, amine optionally substituted by t-butoxycarbonyl; C1 to C12 linear or branched alkyl, aryl, cycloalkyl or heteroalkyl; Z2 is hydrogen, C1 to C12 linear or branched alkyl, aryl, cycloalkyl or heteroalkyl; and B is hydrogen or alkyl.

L73 ANSWER 14 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:867863 CAPLUS Full-text

DOCUMENT NUMBER: 142:71652

TITLE: Process for separating rosmarinic acid from Agastache

rugosa

INVENTOR(S): Cho, Gang Jin; Hwang, Yeong Su; Kim, Jeong Bong; Kim,

Jong Beom; Park, No Dong

PATENT ASSIGNEE(S): Republic of Korea Management : Rural Development

Administration, S. Korea

SOURCE: Repub. Korean Kongkae Taeho Kongbo, No pp. given

CODEN: KRXXA7

DOCUMENT TYPE: Patent LANGUAGE: Korean

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
KR 2001016848	A	20010305	KR 1999-32025	19990804 <
PRIORITY APPLN. INFO.:			KR 1999-32025	19990804 <

AB Provided is a process for separating rosmarinic acid from Agastache rugosa in high purity, wherein the rosmarinic acid is used for producing antioxidizing material and used as additives in food or medical supplies. The process comprises the steps of: drying the Agastache rugosa to have a water content of less than 6 weight and then pulverizing; adding 2-6 times water, based on the weight of the pulverized Agastache rugosa, to obtain the extracted solution of the Agastache rugosa; filtering the extracted solution, performing an adsorption chromatog. and then collecting and condensing activated fractions to obtain the concentrated solution of the rosmarinic acid; performing a gel chromatog. of the concentrated solution and drying fractions containing the rosmarinic acid.

ACCESSION NUMBER: 2004:821008 CAPLUS Full-text

DOCUMENT NUMBER: 141:274556

TITLE: Method for separation and purification of high purity

rosmarinic acid from plant

INVENTOR(S): Kim, Geun Su; Kim, Sam Gon; Kim, Yeong Hoe; Kim, Yong

Ha; Lee, Dong Uk; Lee, Jong Cheol

PATENT ASSIGNEE(S): Korea Ginseng & Tobacco Research Experiment Station,

S. Korea

SOURCE: Repub. Korean Kongkae Taeho Kongbo, No pp. given

CODEN: KRXXA7

DOCUMENT TYPE: Patent LANGUAGE: Korean

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
KR 2001002275	A	20010115	KR 1999-22009	19990614 <
PRIORITY APPLN. INFO.:			KR 1999-22009	19990614 <

AB A method for separating and purifying rosmarinic acid as an antioxidant from a plant, in particular from the leaves of Perilla frutescens var. japonica Hara, using water or a polar solvent is provided; the title method gives highly pure rosmarinic acid at low cost.

=> d 173 ibib abs 60-20

YOU HAVE REQUESTED DATA FROM FILE 'CAPLUS' - CONTINUE? (Y)/N:n

=> d 173 ibib abs 16-20

YOU HAVE REQUESTED DATA FROM FILE 'CAPLUS' - CONTINUE? (Y) /N:y

L73 ANSWER 16 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:799722 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 141:260400

TITLE: An improved process for the preparation of

(+)-(2S,3S)-3-(2-amino-phenylthio)-2-hydroxy-3-(4'-

mehoxyphenyl) propionic acid by using

 $(+)-\alpha$ -methylbenzylamine as a resolving agent INVENTOR(S): Sadanandam, Vennu Sangiah; Shetty, Meera Manjaya;

Sari, Imtiaz Ahmad; Yadav, Jhillu Singh; Rao,

Allavenkata Rama

PATENT ASSIGNEE(S): Council of Scientific & Industrial Research, India

SOURCE: Indian, 14 pp. CODEN: INXXAP

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
IN 182359	A1	19990327	IN 1993-DE1464	19931228 <
PRIORITY APPLN. INFO.:			IN 1993-DE1464	19931228 <
GI				

The invention is directed to an improved process for the preparation of (+)- (2S,3S)-3-(2-amino-phenylthio)-2-hydroxy-3-(4'methoxyphenyl) propionic acid [(+)-(2S,3S)-I], key intermediate in the synthesis of Diltiazem, by using (+)- α -methylbenzylamine (II) as a resolving agent in the presence of a polar solvent and a base. The advantages include high yields of (+)-enantiomer, use of cheap materials and simple methodol. Thus, heating a mixture of (±)- (2RS,3RS)-I and $(+)-\alpha$ -methylbenzylamine in H2O in the presence of LiOH, separating the amine salt [(+)-(2S,3S)-I]•II, and neutralizing it with HCl solution gave acid [(+)-(2S,3S)-I] (m.p. = 136-138°) in 98.77% purity.

L73 ANSWER 17 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:668102 CAPLUS Full-text

DOCUMENT NUMBER: 142:313339

TITLE: Isodon lophanthoides extractant as natural medicine

for treating hepatitis B

INVENTOR(S): Lai, Xiaoping; Hu, Yingjie; Chen, Jiannan; Zhu,

Yutong; Liu, Zhongqiu

PATENT ASSIGNEE(S): Guangzhou University of Traditional Chinese Medicine,

Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 18 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1405177	A	20030326	CN 2002-135069	20021101 <
PRIORITY APPLN. INFO.:			CN 2002-135069	20021101 <

The extractant of Isodon lophanthoides is prepared by extracting with methanol or ethanol under refluxing, then concentrating and adding suitable amount of water, further extracting with petroleum ether, and freeze drying the petroleum ether-insol. phase. The extractant can be prepared by supercrit. CO2 fluid extraction with 2.5-10% ethanol as entrainer at room temperature - 50°. The extractant containing 2-hydroxyursolic acid, 2,19-dihydroxyursolic acid, and rosmarinic acid, can be used for medical formulation (such as tablet, capsule, oral solution, injection, etc.) for treating hepatitis B. Thus, dried Isodon lophanthoides crude powder 4 kg, refluxing in methanol for three times with each time 2 h, after filtration, concentration the solution with reduced pressure, adding water to the residue, then extracted with petroleum ether (60-90°), reducing pressure concentration the petroleum ether insol. part, after freezing dry, crashed the residue and through 80 mesh screen, gave the final Isodon lophanthoides extractant.

L73 ANSWER 18 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:546469 CAPLUS Full-text

DOCUMENT NUMBER: 141:106266

TITLE: Preparation of phenylpropanoic acids derivatives as

selective PPARlpha modulators

INVENTOR(S): Lindstedt Alstermark, Eva-Lotte; Olsson, Anna Christina; Li, Lanna; Aurell, Carl-Johan; Minidis,

Anna; Yousefi-Salakdeh, Esmail; Dahlstrom, Mikael Ulf

Johan

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 43 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
CN, CO, CR,	A1 20040708 AM, AT, AU, AZ, CU, CZ, DE, DK,	WO 2003-GB5602 BA, BB, BG, BR, BW, DM, DZ, EC, EE, EG,	ES, FI, GB, GD,
LK, LR, LS,	LT, LU, LV, MA,	IN, IS, JP, KE, KG, MD, MG, MK, MN, MW, RU, SC, SD, SE, SG,	MX, MZ, NI, NO,
		US, UZ, VC, VN, YU,	
		SD, SL, SZ, TZ, UG,	· · · · · · · · · · · · · · · · · · ·
BY, KG, KZ,	MD, RU, TJ, TM,	AT, BE, BG, CH, CY,	CZ, DE, DK, EE,
ES, FI, FR,	GB, GR, HU, IE,	IT, LU, MC, NL, PT,	RO, SE, SI, SK,
TR, BF, BJ,		GA, GN, GQ, GW, ML,	
CA 2508851		CA 2003-2508851	20031219 <
AU 2003290309	A1 20040714	AU 2003-290309	20031219 <
	B2 20070118		
EP 1572626	A1 20050914		20031219 <
		GB, GR, IT, LI, LU,	
BR 2003017458	A 20051116	CY, AL, TR, BG, CZ, BR 2003-17458	де, но, sk 20031219 <
CN 1753862	A 20060329		20031219 <
CM 1200201	C 20070404		20031219 <
CN 1308291 JP 2006511572	T 20060406		20031219 <
JP 3786945	B2 20060621		20031213
	A1 20050616	US 2004-499893	20040623 <
US 7462644	B2 20081209		
US 20050282822	A1 20051222	US 2004-26806	20041230 <
IN 2005DN02515	A 20070413	IN 2005-DN2515	20050610 <
NO 2005002914	A 20050719	NO 2005-2914	20050615 <
MX 2005PA06812	A 20050908	MX 2005-PA6812	20050621 <
ZA 2005004730	A 20060927	ZA 2005-4730	20050622 <
JP 2006045240	A 20060216	JP 2005-253346	20050901 <
PRIORITY APPLN. INFO.:		GB 2002-29931	A 20021221 <
		SE 2001-4334	A 20011219 <
		WO 2002-GB5738	W 20021218 <
		WO 2002-GB5744	A 20021218 <
		GB 2003-14079	A 20030618
		JP 2004-561668	A3 20031219
		WO 2003-GB5602 WO 2004-EP6597	W 20031219 A 20040617
		US 2005-499261	A2 20050304
OTHER SOURCE(S):	CASREACT 141:10	6266; MARPAT 141:1062	

GΙ

AB Title compds. I [R1 = C1, CF3, CF30; R2 = H, F; R3 = alkyl] and their pharmaceutically acceptable salts, prodrugs were prepared For example, amidation of N-butyl-N-[2-fluoro-4-(trifluoromethyl)benzyl]amine, e.g., prepared from Et (2S)-2-ethoxy-3-(4-hydroxyphenyl)propanoate in 3 steps, and {4-[(2S)-2,3-diethoxy-3-oxopropyl]phenoxy}acetic acid, followed by hydrolysis afforded compound (S)-I [R1 = CF3; R2 = F; R3 = butyl] in 72% yield. Compds. I have EC50 values <0.1 μmol/L for PPARα, e.g., the EC50 value of compound (S)-I [R1 = CF3; R2 = F; R3 = butyl] was 0.001 μmol/L. Of notes, compds. I exhibit improved metabolic stability (in vitro), promising toxicol. profile (no data provided) and particular compds. have the ratio of the EC50 (PPARγ):EC50 (PPARγ) <150:1. Compds. I are claimed useful for the treatment of hypertension, diabetes, etc.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 19 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:546467 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 141:106263

TITLE: Preparation of dimeric dicarboxylic acid derivatives

as PPAR agonists

INVENTOR(S): Sauerberg, Per; Jeppesen, Lone; Polivka, Zdenek;

Sindelar, Karel

PATENT ASSIGNEE(S): Novo Nordisk A/S, Den. SOURCE: PCT Int. Appl., 114 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO.
                        KIND
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                                           APPLICATION NO.
                                                                  DATE
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    WO 2004056740
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                               20040708
                                           WO 2003-DK895
                                                                  20031218 <--
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            CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
            GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
            LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO,
            NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ,
             TM, TN, TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW
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             BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE,
            ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK,
             TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
    US 20040259950
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                                                                  20031212 <--
                         A1
    AU 2003287912
                               20040714
                                           AU 2003-287912
                                                                  20031218 <--
                         A1
    EP 1578716
                                           EP 2003-779752
                                                                  20031218 <--
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    JP 2006510687
                               20060330
                                           JP 2004-561080
                                                                  20031218 <--
PRIORITY APPLN. INFO.:
                                           DK 2002-1966
                                                               A 20021220 <--
                                           US 2003-439410P
                                                             P 20030110
                                           WO 2003-DK895
                                                              W 20031218
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OTHER SOURCE(S): MARPAT 141:106263

GΙ

The title compds. DOC(0) AXLTZUMYBC(0) OE [I; A, B = (un) substituted alkylene, O(alkylene), S(alkylene); D, E = H, alkyl, cycloalkyl; L, M = O, S; T, U = (un) substituted divalent saturated carbon chain, NR1(alkylene) (wherein R1 = H, alkyl); X, Y = (un) substituted arylene, heteroarylene; Z = (un) substituted arylene, heteroarylene, divalent polycyclic ring system] which may be useful in the treatment and/or prevention of conditions mediated by Peroxisome Proliferator-Activated Receptors (PPAR) (no specific biol. data given), were prepared and formulated. E.g., a multi-step synthesis of II, is given. The compds. I are claimed as selective PPARô agonists useful in treating diabetes, syndrome X, cardiovascular diseases, dyslipidemia, and hypercholesteremia.

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS

L73 ANSWER 20 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:453192 CAPLUS Full-text

DOCUMENT NUMBER: 141:6919

TITLE: Preparation of substituted aralkyl derivatives as

antidiabetic, hypolipidemic and hypocholesterolemic

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

agents

INVENTOR(S): Lohray, Braj Bhushan; Lohray, Vidya Bhushan; Jain,

Mukul R.; Basu, Sujay; Pingali, Harikishore; Raval,

Saurin K.; Raval, Preeti S.

PATENT ASSIGNEE(S): Cadila Healthcare Limited, India

SOURCE: PCT Int. Appl., 114 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PA:	TENT NO.			KIN	D	DATE			APPL	ICAT	ION 1	NO.		D	ATE	
WO	20040461	19		A1	_	2004	0603	1	WO 2	003-	IN35	8		2	0031	114 <
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		HR,														
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	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ΤJ,	TM,	TN,
	TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VC,	VN,	YU,	ZA,	ZM,	ZW			
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	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,
	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,
	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD, TG
IN	2002MU00	992		Α		2006	0901		IN 2	002-1	MU99:	2		2	0021	115 <
IN	2003MU00	792		Α			0401								00308	812
CA	2506112			A1		2004	0603		CA 2	003-	2506	112		2	0031	114 <
AU	20033021	11		A1		2004	0615		AU 2	003-	3021	11		2	0031:	114 <
EP	15 6 991 6			A1		2005	0907		EP 2	003-	8083	41		2	0031	114 <
EP	1569916			В1		2009	0107									
	R: AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
	•	SI,	•			RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	SK	
BR	20030157	13		Α		2005	0913		BR 2	003-	1571	3		2	0031	114 <
	1738807			Α		2006	0222								0031	114 <
JP	20065149	7 6		T		2006	0518		JP 2	004-	5703	29		2	0031	114 <
NΖ	540474			Α		2008	0430]	NZ 2	003-	5404	74		2	0031	114 <
MΧ	2005PA05	063		Α		2005	0816]	MX 2	005-1	PA50	63				511 <
ИО	20050024	13		Α		2005	0726]	NO 2	005-	2413				0050	
US	20060142	277		A1		2006	0629	1	US 2	005-	5347	26		2	0051	118 <
CORITY	APPLN.	INFO.	. :						IN 2	002-1	MU99:	2	1	A 2	0021	115 <
									IN 2	003-1	MU79:	2	1	A 2	00308	812

OTHER SOURCE(S): MARPAT 141:6919

GΙ

/ Structure 60 in file .gra /

AB The present invention relates to novel substituted aralkyl derivs. of formula A(CH2)nX-Ar-CH2CH(R)CHR1R2 [A = (substituted) aryl, heteroaryl, heterocyclyl; n = 1-3; X = 0, S; Ar = aromatic, heteroarom. or heterocyclic group; R, R1 = (substituted) amino, (substituted) OH, N3, CN, COOH, tetrazolyl, etc.; R2 = H, alkyl, cycloalkyl], their derivs., their analogs, their tautomeric forms, their pharmaceutically acceptable salts, their pharmaceutically acceptable solvates, pharmaceutical compns. containing them, use of these compds. in medicine and the intermediates involved in their preparation The compds. are useful as antidiabetic, hypolipidemic and hypocholesterolemic agents. Thus, I was prepared, and lowered serum triglyceride in Swiss albino mice by 78%.

=> d 173 ibib abs 21-25 YOU HAVE REQUESTED DATA FROM FILE 'CAPLUS' - CONTINUE? (Y)/N:y

L73 ANSWER 21 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:412810 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 140:423665

TITLE: Preparation of substituted 4-alkoxyoxazole derivatives

as PPAR agonists

INVENTOR(S): Binggeli, Alfred; Grether, Uwe; Hilpert, Hans; Hirth,

Georges; Maerki, Hans-Peter; Meyer, Markus; Mohr,

Peter

PATENT ASSIGNEE(S): F. Hoffmann-La Roche A.-G., Switz.

SOURCE: PCT Int. Appl., 97 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PA.	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION 1	NO.		DATE			
WO	2004	0412	 75		 A1	_	2004	0521		 WO 2	 003-:	 EP12	 189		2	0031	 031 <	(
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	
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OTHER SOURCE(S): CASREACT 140:423665; MARPAT 140:423665
GI
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/ Structure 61 in file .gra /

AB Alkoxyoxazole derivs. of formula I [R1 = alkyl, fluoroalkyl, cycloalkyl, bicyclo-alkyl, tricyclo-alkyl; R2 = H, alkyl, fluoroalkyl; R3-R6 = H, OH, halo, alkyl, etc.; R3R4 = CH=CH-S, alkylene, etc.; R7 = alkyl, fluoroalkyl, aryl, etc.; R8 = H, alkyl] are prepared The compds. are useful for the treatment of diseases such as diabetes. Pharmaceutical compns. containing I are described. Thus, II was prepared in several steps, and had IC50 of 53 nmol/l against PPARα.

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 22 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:308424 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 140:321346

TITLE: Preparation of chiral oxazole-arylpropionic acid

derivatives and their use as PPAR α and PPAR γ agonists for disorders like type II

diabetes

INVENTOR(S): Binggeli, Alfred; Boehringer, Markus; Grether, Uwe;

Hilpert, Hans; Hirth, Georges; Maerki, Hans-Peter;

Meyer, Markus; Mohr, Peter; Ricklin, Fabienne

PATENT ASSIGNEE(S): F. Hoffmann-La Roche A.-G., Switz.

SOURCE: PCT Int. Appl., 108 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004031162	A1	20040415	WO 2003-EP11030	20031006 <
W: AE, AG,	AL, AM, AT	, AU, AZ, E	BA, BB, BG, BR, BY,	BZ, CA, CH, CN,
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GH, GM,	HR, HU, ID	, IL, IN, I	IS, JP, KE, KG, KP,	KR, KZ, LC, LK,
LR, LS,	LT, LU, LV	, MA, MD, M	MG, MK, MN, MW, MX,	MZ, NI, NO, NZ,
OM, PG,	PH, PL, PT	, RO, RU, S	SC, SD, SE, SG, SK,	SL, SY, TJ, TM,

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PRIORITY APPLN. INFO.:
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OTHER SOURCE(S):
                        MARPAT 140:321346
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/ Structure 62 in file .gra /

GT

AΒ The present invention relates to chiral oxazole-arylpropionic acid derivs. (shown as I; variables defined below; e.g. II) and pharmaceutically acceptable salts and esters thereof. The compds. are useful for the treatment and/or prevention of diseases, which are modulated by PPARlpha and/or PPARlpha agonists as e.g. type II diabetes. For I: R1 is aryl or heteroaryl; R2 is H, lower-alkyl, or fluoro-lower-alkyl; R3 and R4 = H, hydroxy, halogen, lower-alkyl, fluorolower-alkyl, hydroxy-lower-alkyl, lower-alkoxy-lower-alkyl, lower-alkoxy, fluoro-lower-alkoxy, hydroxy-lower-alkoxy, lower-alkoxy-lower-alkoxy, or lower-alkenyl, wherein at least one of R3 and R4 is not H; R5 is lower-alkoxy, fluoro-lower-alkoxy, lower-alkenyloxy, fluoro-lower-alkenyloxy, aryloxy, aryllower-alkoxy, or arylfluoro-lower-alkoxy; R6 is H or lower-alkyl; n is 1. EC50 and IC50 values for 10 examples of I towards PPARlpha and PPAR γ are tabulated, e.g. IC50 = 30 and 58 nmol/L for PPAR α and PPAR γ , resp. for II. A method of preparation involving removing a protective ester radical (R6 = protective group) is claimed. Approx. 50 examples prepns. of I are included. For example, II was prepared in 4 steps starting with cyclization of diacetyl monooxime with 4-isopropoxybenzaldehyde to give 2-(4-isopropoxypheny1)-4,5dimethyloxazole 3-oxide hydrochloride, which was converted with POC13 to 4chloromethyl-2-(4-isopropoxyphenyl)-5- methyloxazole, which was coupled to (2S)-2-ethoxy-3-(4-hydroxy-2-methylphenyl)propionic acid Me ester to give (S)-2-ethoxy-3-[4-[2-(4-isopropoxyphenyl)-5-methyloxazol-4-ylmethoxy]-2 $\label{lem:methylphenyl]} \mbox{methylphenyl]} \mbox{propionic acid Me ester, which was hydrolyzed by LiOH to the} \\$

acid.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 23 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:271229 CAPLUS Full-text

DOCUMENT NUMBER: 140:269926

TITLE: Rosmarinic acid-rich Perilla extract, its manufacture

and use as seasoning, and seasoned foods and beverages

INVENTOR(S): Harasawa, Mitsuo; Matsumoto, Katsuyuki; Okutsuka,

Mariko

PATENT ASSIGNEE(S): Ogawa and Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 8 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2004097108	A	20040402	JP 2002-264219	20020910 <
JP 4088127	B2	20080521		
PRIORITY APPLN. INFO.:			JP 2002-264219	20020910 <

AB Title extract is manufactured by extraction of shredded Perilla with alc. solvent at -19 to 5° , then concentration Thus, shredded Perilla frutescens crispa leaves were extracted with aqueous 40% EtOH at -15° for 96 h to give an extract, which had good flavor and contained 0.6% rosmarinic acid.

L73 ANSWER 24 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:220326 CAPLUS Full-text

DOCUMENT NUMBER: 140:270727

TITLE: Preparation of furan derivatives for treatment of

abnormal lipid metabolism, arteriosclerosis, and

diabetes

INVENTOR(S): Hamamura, Kazumasa; Sasaki, Shigekazu; Amano,

Yuichiro; Sakamoto, Junichi; Fukatsu, Kohji

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 325 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT	NO.			KIN	D	DATE		1	APPL	ICAT	ION	NO.		D	ATE	
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JP 2002-261873 A 20020906 <--
JP 2003-185241 A 20030627
WO 2003-JP11308 W 20030904
PRIORITY APPLN. INFO.:
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OTHER SOURCE(S): MARPAT 140:270727

GΙ

/ Structure 63 in file .gra /

AB The title compds. I [wherein R = (un)substituted hydrocarbyl or heterocyclyl; p = 0-2; R1 = H or (un)substituted hydrocarbyl; R2 = (un)substituted aryl; ring A = (un)substituted aromatic ring; X1 = 0 or S; X2 = a bond, O, S, SO, or SO2; Y = a bond, O, S, SO, SO2, CO, (un)substituted CONH, or NHCO; M1-M3 = independently a bond or (un)substituted aliphatic hydrocarbyl; M4 = (un)substituted aliphatic hydrocarbyl; with exclusions], or prodrugs, or pharmaceutically acceptable salts thereof are prepared For example, the compound II was prepared in a multi-step synthesis. II exhibited EC50 of 0.10 μM towards human G protein-coupled receptors (GPR40). I are useful for the treatment of abnormal lipid metabolism, arteriosclerotic diseases, secondary diseases, diabetes, etc. (no data). Formulations containing I as an active ingredient were also described.

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 25 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:203818 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 140:253565

TITLE: Preparation of novel 2-arylthiazole compounds as

peroxisome proliferator activated receptors

(PPARlpha and PPAR γ) agonists

INVENTOR(S): Binggeli, Alfred; Grether, Uwe; Hilpert, Hans; Hirth,

Georges; Maerki, Hans-Peter; Meyer, Markus; Mohr,

Peter

PATENT ASSIGNEE(S): F.Hoffmann-La Roche A.-G., Switz.

SOURCE: PCT Int. Appl., 169 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT	NO.			KIN	D	DATE			APPL	ICAT	I NOI	NO.		D	ATE	
WO 2004	0204	20		A1	_	2004	0311		WO 2	003-	EP92	81		2	0030	821 <
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PRIORITY APPLN. INFO.:
                                            EP 2002-19146
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                                            WO 2003-EP9281
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OTHER SOURCE(S): MARPAT 140:253565

GI

/ Structure 64 in file .gra /

The present invention relates to compds. of formula (I) [R1 = aryl, AB heteroaryl; R2 = H, lower-alkyl, fluoro-lower-alkyl; R3, R4, R5 = H, H0, lower-alkenyl, halogen, lower-alkyl, fluoro-lower-alkyl, hydroxy-lower-alkyl, lower-alkoxy-lower- alkyl, lower-alkoxy, fluoro-lower-alkoxy, hydroxy-loweralkoxy, lower-alkoxy-lower-alkoxy; wherein at least one of R3, R4, R5 and R6 is not hydrogen, or R3 and R4 are bonded to each other to form a ring together with the carbon atoms to which they are attached, and R3 and R4 together are -CH:CH-S-, -S-CH:CH-, -CH:CH-O-, -O-CH:CH-, -CH;CHCH:CH-, -(CH2)3-5, -O-(CH2)2-3- or -(CH2)2-3-0-, and R5 and R6 are as defined above; R7 = lower-alkyl, lower-alkoxy, lower-alkenyloxy, aryloxy, aryl-lower-alkoxy; R5 = H, loweralkyl; R9, R10 = H, lower-alkyl, lower-alkenyl, cycloalkyl, Ph, -[1,3]dioxan-2-ethyl; n = 1-3] and pharmaceutically acceptable salts and/or pharmaceutically acceptable esters thereof. The compds. I exceed the compds. known in the art, in as much as they bind to and activate both, PPARlpha and PPARy, simultaneously and very efficiently. Therefore, these compds. I combine the anti-qlycemic effect of PPARy activation with the anti-dyslipidemic effect of PPARlpha activation. Consequently, plasma glucose and insulin are reduced (i.e. insulin sensitization), triglycerides lowered and HDL cholesterol increased (i.e. improved lipid profile) and in addition, such compds. may also lower LDL cholesterol, decrease blood pressure, and counteract inflammatory atherosclerosis. Thereby, the compds. I are useful for the treatment of diseases such as diabetes, noninsulin dependent diabetes mellitus, elevated blood pressure, increased lipid and cholesterol levels, atherosclerotic diseases, metabolic syndrome, endothelial dysfunction, procoagulant state, dyslipidemia, polycystic ovary syndrome, inflammatory diseases, or proliferative diseases. For example, rac-3-[4-[3-[2-(4-tert-butylphenyl)-5methylthiazol-4-yl]propoxy]-2- methylphenyl]-2-ethoxypropionic acid showed IC50 of 15 and 20 nM against PPAR α and PPAR γ , resp., in luciferase transcriptional reporter gene assays using baby hamster kidney cells transfected with either pFA-PPAR γ -LBD or pFA-PPARlpha-LBD expression plasmids and a reporter plasmid.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d 173 ibib abs 30-40 YOU HAVE REQUESTED DATA FROM FILE 'CAPLUS' - CONTINUE? (Y)/N:y

L73 ANSWER 30 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:826813 CAPLUS Full-text

DOCUMENT NUMBER: 139:311997

TITLE: Rosmarinic acid compositions containing fragrances or

cationic surfactants

INVENTOR(S): Someya, Keita; Mizushima, Yukako; Matsukawa, Hiroshi

PATENT ASSIGNEE(S): Lion Corp., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 140 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATI	ENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2	2003300811	A	20031021	JP 2002-104240	20020405 <
JP :	3821222	B2	20060913		
JP 2	2006249092	A	20060921	JP 2006-90601	20060329 <
PRIORITY	APPLN. INFO.:			JP 2002-104240 A	3 20020405 <

The compns., useful for flavoring materials, perfumes, and hair or fiber treatment agents, contain rosmarinic acid (I) and fragrances or cationic surfactants. I improves fragrance stability and hand feel of hair and fiber products. A fragrance composition comprising fragrant preparation 0.01, I 0.1, 1,3-butanediol 30, and H2O to 100 weight% showed good fragrance after storage in a cup at room temperature for 1 day.

L73 ANSWER 31 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:796655 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 139:292053

TITLE: Etherification process for the preparation of

2-ethoxy-3-[4-[2-(4-

methanesulfonyloxyphenyl)ethoxy]phenyl]propanoic acid

derivatives

INVENTOR(S): Larsson, Maria

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 9 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAI	ENT	NO.			KIN	D	DATE		i	APPL	ICAT	ION	NO.		D	ATE		
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WO	2003	0828	12		A2		2003	1009	Ţ	WO 2	003-0	GB13	95		20	00303	328	<
WO	2003	0828	12		А3		20040108											
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PRIORITY APPLN. INFO.:
                                            SE 2002-1005
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OTHER SOURCE(S):
                   CASREACT 139:292053; MARPAT 139:292053
GΙ
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

An efficient industrial-scale process for the preparation of 2-ethoxy-3-[4-[2-(4-methanesulfonyloxyphenyl)ethoxy]phenyl]propanoic acid derivs. [I; R = H, acid-protecting group; 1-(S)-2-ethoxy-3-[4-[2-(4-methanesulfonyloxyphenyl)ethoxy]phenyl]propanoic acid] is described which comprises the etherification of 2-ethoxy-3-(4-hydroxyphenyl)propanoate derivs. [II; e.g., Et <math>(S)-2-ethoxy-3-(4-hydroxyphenyl)propanoate] with 2-(4-methanesulfonyloxyphenyl)ethyl derivs. [III; X = leaving group; e.g., 2-(4-methanesulfonyloxyphenyl)ethyl methanesulfonate] in the presence of a base (e.g., sodium carbonate) and using water as a diluent.

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 32 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:696734 CAPLUS Full-text DOCUMENT NUMBER: 139:230768

TITLE: Preparation of (arylalkyl)thiazoles and oxazoles as peroxisome proliferator activated receptor modulators

for treating diabetes mellitus, syndrome X, and

cardiovascular disease

INVENTOR(S): Conner, Scott Eugene; Knobelsdorf, James Allen;

Mantlo, Nathan Bryan; Schkeryantz, Jeffrey Michael; Shen, Quanrong; Warshawsky, Alan M.; Zhu, Guoxin

Shell, Qualifolig, warshawsky, Afan Fr., 2

PATENT ASSIGNEE(S): Eli Lilly and Company, USA SOURCE: PCT Int. Appl., 223 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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APPLICATION NO.
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            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
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PRIORITY APPLN. INFO.:
                                         WO 2003-US2679
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OTHER SOURCE(S): MARPAT 139:230768
GΙ
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/ Structure 65 in file .gra /

Title compds. I [wherein R3, R4, R30, and R40= independently H, alkyl, halo, AΒ or alkoxy; R5 = (un)substituted alkyl, alkenyl, aryl(oxy)alkyl, or arylthioalkyl; or when R5 = alkyl, R5 may be combined with <math>W to form a heterocycloalkyl fused to the oxazole or thiazole ring; R6 = trihalomethyl, trihalomethoxy, (hydroxy)alkyl, alkylcarbamoyl, tetramethyldioxaborolanyl, halo, alkanoyl, carboxyalkoxy, (cyclo)alkoxy, tetrahydropyranyloxy, morpholinyl, or (un)substituted aryloxy, arylthio, heterocyclyloxy, pyridinyl, pyrimidinyl, pyrazinyl, or arylalkyl; R7 and R8 = independently H, CF3, or alkyl; R9 = (un)substituted (aryl)alkyl or alkenyl; R10 = H or alkyl; Q = a bond, O, or CH2; T1 = C or N; W = CH2, O, OCH2, S, SO2, or (un)substituted CONH, NH, or NHCH2; X = C, CH2C, or CCH2; Y and Z = independently O, N, or S wherein at least 1 of Y and Z = O or S; A = CO2H, alkylnitrile, CONH2, or (CH2)nCO2R19; n = 0-3; R19 = H or alkyl; and pharmaceutically acceptable salts thereof] were prepared as peroxisome proliferator activated receptor δ (PPAR δ) modulators (no data). For example, (4-mercapto-2-methylphenoxy) acetic acid Et ester was condensed with 1-[4-[2-(2-chloro-6-fluorophenyl)]-2-(4trifluoromethylphenyl)thiazol-5-yl]ethanol in the presence of PBu3 and 1,1'-(azodicarbonyl) bipiperidine in toluene. Deesterification with LiOH in THF produced II. I and their pharmaceutical compns. are useful for the prevention and or treatment of diabetes mellitus, syndrome X, and cardiovascular disease (no data).

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 33 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:678776 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 139:214119

TITLE: Preparation of 3-phenylpropionic acid derivatives for

treatment of diabetes and hyperlipemia

INVENTOR(S): Kawanishi, Masashi; Umeno, Hiroshi PATENT ASSIGNEE(S): Asahi Kasei Kabushiki Kaisha, Japan

SOURCE: PCT Int. Appl., 546 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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AU	2003	2113	83		A1		2003	0909		AU 2	003-	2113	83		2	0030	218	<
EP	1484	316			A1		2004	1208		EP 2	003-	7052	66		2	0030	218	<
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CN	1296	349			С		2007	0124										
US	2004	0072	690		A1		2004	0415		US 2	003-	3678	57		2	0030	219	<
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										WO 2	003-	JP16	95		W 2	0030	218	
THER SO	TIRCE	(S) ·			MARI	РΔТ	139.	2141	19									

OTHER SOURCE(S): MARPAT 139:214119

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/ Structure 66 in file .gra /

AB The title compds. I [wherein R1 = (un)substituted alkyl or Ph; R2 = (un)substituted alkyl with exclusions; R3 = halo, alkyl, or alkoxy; m = 0-4; R4 = alkyl; R5 = H or alkyl; n = 2-4; X = NH or O] and salts thereof are prepared I are highly effective in lowering blood sugar, lipid, and total cholesterol, and are useful for the treatment of diabetes and hyperlipemia. For example, compound II was prepared in a multi-step synthesis. II lowered

42% of blood sugar and 51% of lipids in rat in the amount of 1 mg/kg in 15 days.

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 34 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:633683 CAPLUS Full-text

DOCUMENT NUMBER: 139:185673

TITLE: Preparation and compositions of polymorphic forms of

bicyclic antidiabetic agents

INVENTOR(S): Srisilla, Raju; Potlapally, Rajender Kumar;

Mamillapalli, Ramabhadra Sarma; Gaddam, Om Reddy

PATENT ASSIGNEE(S): Reddy's Laboratories Limited, India

SOURCE: PCT Int. Appl., 35 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	PATENT NO.						DATE		•	APPL:	ICAT:	ION 1	. O <i>I</i>		D	ATE	
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IN	2002	MA00	095		A		2005	0304		IN 2	002-1	MA95			2	0020	207 <
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AB This invention relates to novel polymorphic/pseudopolymorphic forms and compns. of arginine salt of 3-[4-[2-(3,4-dihydro-1,4-benzothiazin-4-yl)ethoxy]phenyl]-2-ethoxypropanoic acid, preferably, L-arginine salt of (2S)-3-[4-[2-(3,4-dihydro-1,4-benzothiazin-4-yl)ethoxy]phenyl]-2- ethoxypropanoic acid. The polymorphic forms of the present invention are more active, as antidiabetic and hypolipidemic agent, than the 3-[4-[2-(2,3-dihydro-1,4-benzothiazin-4-yl)ethoxy]phenyl]-2- ethoxypropanoic acid.

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 35 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:570941 CAPLUS Full-text

DOCUMENT NUMBER: 139:133829

TITLE: Processes for the preparation of glutamic acid

compounds and intermediates thereof and novel

intermediates used in the processes

INVENTOR(S): Kawahara, Shigeru; Amino, Yusuke; Mori, Kenichi;

Funakoshi, Nao; Takemoto, Tadashi

PATENT ASSIGNEE(S): Ajinomoto Co., Inc., Japan

SOURCE: PCT Int. Appl., 95 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

PA'	TENT	NO.			KIN		DATE			APPL	ICAT	ION 1	NO.		Γ	ATE		
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RU	2305	677			C2		2007	0910		RU 2	004-	1229	10		2	20021	129	<
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										US 2	004-	8725	73		A3 2	20040	622	
										US 2	005-	2839	43		A3 2	20051	122	
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OTHER SOURCE(S): CASREACT 139:133829; MARPAT 139:133829

AB This document discloses the following: a process for preparing industrially and efficiently glutamic acid compds. (such as monatin) useful as sweeteners or intermediates for the production of drugs or the like; a process for preparation of intermediates used therein; novel intermediates included among them; a process for preparation of optically active monatin; a process for preparation of intermediates used therein; and novel intermediates included among them. Specifically, this document discloses a process for the preparation of glutamic acid compds. (or salts thereof) including monatin, which comprises preparing a ketoglutaric acid compound serving as a precursor of the target glutamic acid compound by condensing a specific pyruvic acid compound with oxalacetic acid or pyruvic acid through cross-aldol reaction and, if necessary, subjecting the obtained condensate to decarboxylation, and replacing the carbonyl group of the ketoglutaric acid compound by an amino group.

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 36 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:511336 CAPLUS Full-text DOCUMENT NUMBER: 139:85372

TITLE: Preparation of pyrazolopyrimidines and related

compounds as hPPARa and hPPARy ligands

INVENTOR(S): Das, Saibal Kumar; Bhuniya, Debnath; Madhavan, Gurram

Ranga; Iqbal, Javed; Chakrabarti, Ranjan

Reddy's Laboratories Ltd., India

SOURCE: PCT Int. Appl., 139 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT ASSIGNEE(S):

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											WO 2	002-	IB54	42		W 2	0021	217 <
OTHER GI	SOU	JRCE	(S):			CAS:	REAC	T 13	9:85	372;	MAR	PAT	139:	8537.	2			

/ Structure 67 in file .gra /

AΒ Title compds. I [R1 = H, halo, OH, etc.; R2 = H, OH, halo, etc.; R3 = H, (un) substituted alkyl, cycloalkyl, etc.; Z = O, NR4; R4 = H, (un) substituted alkyl, aryl, etc.; Y = O, S, NR6, etc.; R6 = H, (un)substituted alkyl, aryl, etc.; Ar = (un)substituted aromatic, heteroarom., heterocyclic; G = O, S; X = O, NHR5, (CH2)pO, etc.; R5 = H, (un)substituted alkyl, aryl, etc.; n = 1-4; p = 0-4; A = (un)substituted pyrazolopyrimidine, imidazolopyrimidine] and their pharmaceutically acceptable salts and formulations were prepared For example, O-alkylation of 5-ethyl-1,4-dihydro-1-methyl-3-propyl-7H-pyrazolo[4,3d]pyrimidin-7-one by chloroacetyl II, e.g., prepared from 4-aminothiophenol in 3-steps, followed by ester hydrolysis, afforded claimed pyrazolopyrimidine III in 5% yield. In hPPAR α and hPPAR γ Luciferase ligand binding assays, 2examples of compds. I, e.g., pyrazolopyrimidine III, exhibited activity at 50 and 1 μM , resp. The test compds. also inhibited HMG CoA reductase (no data provided). Compds. I are claimed useful as antidiabetic, hypolipidemic, antiobesity and hypocholesterolemic agents.

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 37 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:504661 CAPLUS Full-text DOCUMENT NUMBER: 139:68323

TITLE: Manufacture of extracts containing phenols from salted

red Perilla frutescens leaves

INVENTOR(S): Natsume, Midori; Osakabe, Naoko; Kashiwazaki, Hideaki

PATENT ASSIGNEE(S): Meiji Seika Kaisha, Ltd., Japan; Toyotama Perfumery

Co., Ltd.

SOURCE: Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2003180286	A	20030702	JP 2001-388962	20011221 <
JP 3775584	В2	20060517		

PRIORITY APPLN. INFO.:

JP 2001-388962 20011221 <--

AB The exts. containing phenols are manufactured by extraction of components from salted red leaves of Perilla frutescens with hydrophilic solvents under acidic and heating conditions, chromatog. of the exts. with adsorptive resins, and concentration and pulverization of the exts. The exts. containing antiallergy phenols (e.g., rosmarinic acid) are useful for foods and beverages for treatment of pollinosis, atopic dermatitis, etc.

L73 ANSWER 38 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:491169 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 139:69054

TITLE: Preparation of substituted phenylpropionic acid

derivatives as agonists to human peroxisome proliferator-activated receptor alpha (PPAR) Lindstedt Alstermark, Eva-Lotte: Olsson, Anna

INVENTOR(S): Lindstedt Alstermark, Eva-Lotte; Olsson, Anna

Christina; Li, Lanna

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 43 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PAT	[ENT	ΝΟ.			KIN	D	DATE			APPL	ICAT	ION 1	ΝΟ.		D	ATE	
WO	2003	0518	22		A1	_	2003	0626		WO 2	002-	GB57	44		2	0021	218 <
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                                         WO 2004-EP6597
                                                            A 20040617
                                         US 2005-499261 A2 20050304
                      MARPAT 139:69054
OTHER SOURCE(S):
```

/ Structure 68 in file .gra /

GI

AB The present invention provides the S enantiomer of a compound of formula (I) (wherein R1 represents 2,4-difluorophenyl or cyclohexyl) as well as pharmaceutically acceptable salts, solvates, crystalline forms and prodrugs thereof, processes for preparing such compds., their the utility in treating clin. conditions including lipid disorders (dyslipidemias) whether or not associated with insulin resistance, methods for their therapeutic use, and pharmaceutical compns. containing them. Thus, to a solution of [4-((2S)-2,3diethoxy-3-oxopropyl)phenoxylacetic acid (0.108 g) 3.6 mL CH2Cl2 were added N-(cyclohexylmethyl)-N-heptylamine hydrochloride (0.090 g) and DMAP (0.098 g) followed by 1-ethyl-3-(3- dimethylaminopropyl)carbodiimide hydrochloride (0.070 g) and the reaction mixture was stirred at room temperature overnight to give, after workup and silica gel chromatog., Et (2S)-3-[4-[2-[(cyclohexylmethyl)(heptyl)amino]-2- oxoethoxy]phenyl]-2-ethoxypropanoate which (0.031 g) was saponified with LiOH in aqueous THF at room temperature overnight and acidified with aqueous 2 M HCl to give (2S)-3-[4-[2-[(cyclohexylmethyl)(heptyl)amino]-2-oxoethoxy]phenyl]-2- ethoxypropanoic acid. The compds. I had EC50 of less than $0.5~\mu mol/L$ for PPAR α and preferred compds. have EC50 of less than 0.05 μ mol/L for PPAR α . They were more potent with respect to PPAR α than with respect to PPAR γ .

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 39 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:491168 CAPLUS Full-text

DOCUMENT NUMBER: 139:69049

TITLE: Preparation of substituted phenylpropionic acid

derivatives as agonists to human peroxisome proliferator-activated receptor alpha (PPAR)

INVENTOR(S): Lindstedt Alstermark, Eva-Lotte; Olsson, Anna

Christina; Li, Lanna

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

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ΑT	3387	43			T		2006	0915		AT 2	002-	8049	64			0021		
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JP 2006-123399 20060427 <--
SE 2001-4334 A 20011219 <--
CN 2002-828123 A3 20021218 <--
JP 2003-552709 A3 20021218 <--
JP 2003-552710 A3 20021218 <--
WO 2002-GB5738 W 20021218 <--
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WO 2003-GB5602 A 20031219
WO 2004-EP6597 A 20040617
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PRIORITY APPLN. INFO.:
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OTHER SOURCE(S): MARPAT 139:69049

GΙ

/ Structure 69 in file .gra /

The S enantiomer of I, n = 1 or 2, (C6H13 = hexyl) as well as their AΒ pharmaceutically acceptable salts, solvates, crystalline forms and prodrugs are synthesized using various solvents and in presence of charcoal-supported palladium catalyst. The utility of these compds. in clin. conditions such as lipid disorders (dyslipidemias) whether or not associated with insulin resistance and therapeutic and other pharmaceutical activities is also investigated. For example, (2S)-3-(4{2-[benzyl(hexyl)amino]-2oxoethoxy}phenyl)2-ethoxypropionic acid was prepared in 58% yield via reaction of (2S)-2-ethoxy-3-(4-hydroxyphenyl)propanoate and benzyl bromoacetate. THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L73 ANSWER 40 OF 600 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:454296 CAPLUS Full-text

DOCUMENT NUMBER: 139:36527

Preparation of imidazolidinone derivatives as TITLE:

> peroxisome proliferator activated receptor agonists Gibson, Tracey Ann; Johnston, Richard Duane; Mantlo, Nathan Bryan; Thompson, Richard Craig; Wang, Xiaodong;

Winneroski, Leonard Larry, Jr.; Xu, Yanping

PATENT ASSIGNEE(S): Eli Lilly and Company, USA SOURCE: PCT Int. Appl., 408 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Pat.ent. LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

INVENTOR(S):

PATENT NO.	KIND DATE	E APPLIC	CATION NO.	DATE
WO 2003048130	A2 2003	30612 WO 200	2-US36128	20021126 <
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OTHER SOURCE(S):
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OTHER SOURCE(S): MARPAI 139:3632/

GΙ

/ Structure 70 in file .gra /

AΒ The present invention is directed to compds. represented by the following structural Formula (I) [wherein R1 = H, each (un) substituted C1-C8 alkyl, ary1-C0-4-alky1, heteroary1-C0-4-alky1, C3-6 cycloalky1ary1-C0-2-alky1, or CH2-C(0)-R17-R18 (wherein R17 = 0, NH; R18 = optionally substituted benzyl); R2 = C1-6 alkyl, C1-6 alkenyl, aryl-C0-4-alkyl, heteroaryl-C0-4-alkyl, C1-4 alkylsulfonamide, C1-4 alkylamide, OH, C1-4 alkoxy, C3-6 cycloalkyl; W = O, S; X =an optionally substituted C1-5 alkylene linker wherein one carbon atom of the linker may optionally be replaced with O, NH, S, and optionally two carbons together may form a double bond; Y = C, O, S, NH, a single bond; E =C(R3)(R4)A, A, (CH2)nCO2R19; wherein A = CO2H, C1-3 alkylnitrile, carboxamide, each (un) substituted sulfonamide, acylsulfonamide, tetrazole, or isoxazole; R3 = H, C1-5 alkyl, C1-5 alkoxy; R4 = H, halo, each (un)substituted C1-5 alkyl, C1-5 alkoxy, C3-6 cycloalkyl, aryl-C0-4-alkyl, aryl-C0-2 alkoxy, or Ph; or R3 and R4 are combined to form a C3-8 cycloalkyl; R19 = H, each (un)substituted arylmethyl or C1-4 alkyl; n = 0-3; R21 = H, oxo, each (un)substituted C1-6alkyl, aryl, C1-4 alkylaryl, or heteroaryl; R22 = H, each (un)substituted C1-6 alkyl, aryl, C1-4 alkyl-aryl, or heteroaryl]. These compds. are useful for preventing or treating diabetes mellitus or treating syndrome X or cardiovascular disease (no data). Thus, To a solution of 2-methyl-2-[2methyl-4-[2-(3-methyl-2-oxoimidazolidin-4-yl)ethoxy] phenoxy] propionic acid Et ester (0.040 g) in DMF (2.0 mL), was added NaH (60% in mineral oil, 0.0066 g)in one portion and the mixture was stirred for 15 min at room temperature, treated with 4-tert-butylbenzyl bromide (0.030 mL), and stirred for 4 h at room temperature to give, after workup, an Et ester intermediate, which was treated with a mixture of MeOH (2 mL)/5.0 N NaOH (1 mL) at room temperature overnight, concentrated, diluted with water (2 mL), cooled down to 0°, and acidified to pH 2 by adding concentrated HCl dropwise to give, after purification on a Chem elut 1005 tube, 2-[4-[2-[1-(4-tert-Butylbenzyl)-3methyl-2-oxoimidazolidin-4-yl]ethoxy]-2- methylphenoxy]-2-methylpropionic acid

as an colorless oil (0.022 g, 42%).

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s 11 sss sam SAMPLE SEARCH INITIATED 12:49:53 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 2561 TO ITERATE

78.1% PROCESSED 2000 ITERATIONS 50 ANSWERS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE**

PROJECTED ITERATIONS: 48185 TO 54255 PROJECTED ANSWERS: 3861 TO 5717

50 SEA SSS SAM L1 1.2

=> d scan

REGISTRY COPYRIGHT 2009 ACS on STN L250 ANSWERS

ΙN Benzenepropanoic acid, α , 2-diethoxy-4-[2-(5-methyl-2-phenyl-4thiazolyl)ethoxy]-, ethyl ester

MF C27 H33 N O5 S

/ Structure 84 in file .gra /

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

50 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN 1.2

Benzenepropanoic acid, α -ethoxy-4-[2-(4-ethoxy-9H-carbazol-9-ΤN yl)ethoxy]-, ethyl ester

C29 H33 N O5 MF

=> s l1 sss full

FULL SEARCH INITIATED 12:51:33 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 50873 TO ITERATE

100.0% PROCESSED 50873 ITERATIONS 4362 ANSWERS

SEARCH TIME: 00.00.01

L3 4362 SEA SSS FUL L1

=> s 13/prep

1060 L3

4706241 PREP/RL

821 L3/PREP T.4

(L3 (L) PREP/RL)

=> s 14 and (py<2003 or ay<2003 or pry<2003) 22983071 PY<2003

4502933 AY<2003 3971676 PRY<2003

L5 587 L4 AND (PY<2003 OR AY<2003 OR PRY<2003)

=> S L5 AND HYDROGENA?

301729 HYDROGENA?

L6 49 L5 AND HYDROGENA?

=> S L6 AND CHIRAL?

139312 CHIRAL?

L7 6 L6 AND CHIRAL?

=> S L6 AND (CHIRAL? OR OPTICAL?)

139312 CHIRAL? 1145559 OPTICAL?

L8 8 L6 AND (CHIRAL? OR OPTICAL?)

=> D L8 1-8 IBIB ABS TI HIT

L8 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2007:544712 CAPLUS Full-text

DOCUMENT NUMBER: 148:449648

TITLE: An improved process for the preparation of phenoxazine

antidiabetic compounds

INVENTOR(S): Rao, Siripragada Mahender; Reddy, Chepyala Naveen

Kumar; Reddy, Challa Maheedhara; Sarma, Mamillapalli

Ramabhandra; Reddy, Gaddam Om

PATENT ASSIGNEE(S): Reddy's Laboratories Ltd., India

SOURCE: Indian Pat. Appl., 18pp.

CODEN: INXXBQ

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

-----IN 2002MA00508 A 20070511 IN 2002-MA508 20020708 <-
PRIORITY APPLN. INFO.: IN 2002-MA508 20020708 <--

OTHER SOURCE(S): CASREACT 148:449648; MARPAT 148:449648

GΙ

/ Structure 99 in file .gra /

The invention relates to an improved process for the preparation of antidiabetic phenoxazinylethoxyphenylalkoxypropanoate arginine salts I [R1 = Me, Et, Pr, i-Pr]. The process uses inexpensive chems, and an easy resolution via chiral amine salts. Thus, condensation of 4-[2-(phenoxazin-10-yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the corresponding glycidic ester, which was hydrogenated over Pd/C, O-ethylated with di-Et sulfate in xylene, and saponified with aqueous NaOH in MeOH to give acid (±)-II. Resolution of this racemate using (-)-ephedrine and salification with L-arginine in i-PrOH gave I (R1 = Et).

TI An improved process for the preparation of phenoxazine antidiabetic

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compounds
    PATENT NO.
                   KIND DATE APPLICATION NO. DATE
                      ----
                                         ______
                       A 20070511 IN 2002-MA508
    IN 2002MA00508
                                                               20020708 <--
                             20020708 <--
PRAI IN 2002-MA508
     The invention relates to an improved process for the preparation of
     antidiabetic phenoxazinylethoxyphenylalkoxypropanoate arginine salts I [R1 =
     Me, Et, Pr, i-Pr]. The process uses inexpensive chems. and an easy resolution
     via chiral amine salts. Thus, condensation of 4-[2-(phenoxazin-10-
     yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the
     corresponding glycidic ester, which was hydrogenated over Pd/C, O-ethylated
     with di-Et sulfate in xylene, and saponified with aqueous NaOH in MeOH to give
     acid (±)-II. Resolution of this racemate using (-)-ephedrine and salification
     with L-arginine in i-PrOH gave I (R1 = Et).
    222834-12-0F 222834-14-2F 222834-21-1F 222834-30-2F
ΤT
    267228-45-5P 493014-69-0P 493014-71-4P 493014-73-6P 493014-77-0P
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    493014-84-9P 493014-96-3P 493014-97-4P 493014-98-5P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP
    (Preparation); RACT (Reactant or reagent)
        (improved preparation of phenoxazine antidiabetic compds.)
    ANSWER 2 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2003:76765 CAPLUS <u>Full-text</u>
                      138:137318
DOCUMENT NUMBER:
                       An improved process for the preparation of
TITLE:
                       antidiabetic phenoxazine compounds, e.g., ragaglitazar
                        L-arginine salt
                        Siripragada, Mahender Rao; Chepyala, Naveen Kumar
INVENTOR(S):
                        Reddy; Challa, Maheedharareddy; Mamillapalli,
                        Ramabhadra Sarma; Gaddam, Om Reddy
PATENT ASSIGNEE(S):
                        Reddy's Laboratories Ltd., India
SOURCE:
                       PCT Int. Appl., 31 pp.
                       CODEN: PIXXD2
DOCUMENT TYPE:
                       Patent
                       English
LANGUAGE:
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
    PATENT NO.
                 KIND DATE APPLICATION NO. DATE
    WO 2003008397 A1 20030130 WO 2002-IB2776 20020716 <--
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            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
            PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
            UA, UG, US, UZ, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
            CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
            PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
            NE, SN, TD, TG
                       A1
                             20031220
                                          IN 2001-MA585
                                                                20010718 <--
    IN 191728
    IN 2001MA00585 A 20081128
AU 2002313569 A1 20030303
                                         AU 2002-313569 20020716 <--
IN 2001-MA585 A 20010718 <--
WO 2002-IB2776 W 20020716 <--
PRIORITY APPLN. INFO.:
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OTHER SOURCE(S): CASREACT 138:137318; MARPAT 138:137318

AΒ The invention relates to an improved, multi-step process for the preparation of antidiabetic compds. I [where R1 represents an alkyl group such as Me, Et, Pr, iso-Pr, Bu, tert-Bu, and the like]. I are known antidiabetic, hypolipidemic, and antihypercholesterolemic agents. Older methods of preparing I are industrially unsuitable for a variety of reasons, including exotic and moisture-sensitive reactions, tedious purifns., low yields, and long cycle times. The new method is simple, robust, scalable, and provides I with high chemical and chiral purity. The use of highly reactive, difficultto-handle, and expensive chems. is avoided, and simple, inexpensive chems. such as di-Et sulfate and K2CO3 are used instead. Two variants of the process are claimed and illustrated, in which the resolution step occurs at different points. For instance, Darzen's condensation of 4-[2-(phenoxazin-10yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the corresponding glycidic ester, which was hydrogenated over Pd/C and saponified with aqueous NaOH in MeOH to give acid (±)-II. Two-step resolution of this racemate using (S)-phenylglycinol and then (-)- α -methylbenzylamine gave (S)-II of 99.2% purity. Combined etherification/esterification of the latter with NaH and EtI, saponification of the ester with aqueous NaOH in MeOH, and salification with L-arginine in iso-PrOH gave I (R1 = Et), i.e., the arginine salt of ragaglitazar.. The alternative sequence involves Darzen's condensation, hydrogenation, etherification/esterification, ester saponification, resolution, and salification.

TI An improved process for the preparation of antidiabetic phenoxazine compounds, e.g., ragaglitazar L-arginine salt

REFERENCE COUNT: THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT KIND DATE APPLICATION NO. DATE PATENT NO. 20030130 WO 2002-IB2776 WO 2003008397 A1 20020716 <--W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG 20031220 IN 2001-MA585 20010718 <--IN 191728 Α1 IN 2001MA00585 20081128 Α AU 2002313569 A1 20030303 AU 2002-313569 20020716 <--PRAI IN 2001-MA585 20010718 <--Α WO 2002-IB2776 \mathbb{W} 20020716 <--

The invention relates to an improved, multi-step process for the preparation of antidiabetic compds. I [where R1 represents an alkyl group such as Me, Et, Pr, iso-Pr, Bu, tert-Bu, and the like]. I are known antidiabetic, hypolipidemic, and antihypercholesterolemic agents. Older methods of preparing I are industrially unsuitable for a variety of reasons, including exotic and moisture-sensitive reactions, tedious purifns., low yields, and long cycle times. The new method is simple, robust, scalable, and provides I with high chemical and chiral purity. The use of highly reactive, difficult-to-handle, and expensive chems. is avoided, and simple, inexpensive chems. such as di-Et sulfate and K2CO3 are used instead. Two variants of the process

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are claimed and illustrated, in which the resolution step occurs at different
points. For instance, Darzen's condensation of 4-[2-(phenoxazin-10-
yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the
corresponding glycidic ester, which was hydrogenated over Pd/C and saponified
with aqueous NaOH in MeOH to give acid (±)-II. Two-step resolution of this
racemate using (S)-phenylglycinol and then (-)-\alpha-methylbenzylamine gave (S)-II
of 99.2% purity. Combined etherification/esterification of the latter with
NaH and EtI, saponification of the ester with aqueous NaOH in MeOH, and
salification with L-arginine in iso-PrOH gave I (R1 = Et), i.e., the arginine
salt of ragaglitazar.. The alternative sequence involves Darzen's
condensation, hydrogenation, etherification/esterification, ester
saponification, resolution, and salification.
222834-12-0F, Ethyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2-
ethoxypropanoate 222834-14-2P, Ethyl
3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-hydroxypropanoate
222834-21-1P, 3-[4-[2-(Phenoxazin-10-yl)ethoxy]phenyl]-2-ethoxypropanoic
      222834-23-3P, 3-[4-[2-(Phenoxazin-10-yl)ethoxy]phenyl]-2-
                       222834-30-2P,
hydroxypropanoic acid
(S)-3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-ethoxypropanoic acid
493014-69-0P, Ethyl 3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-2,3-
epoxypropanoate
                 493031-09-7P 493031-10-0P
RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic
preparation); PREP (Preparation); RACT (Reactant or reagent)
   (intermediate; improved preparation of antidiabetic phenoxazine derivs.,
   e.g., ragaglitazar arginine salt)
222834-24-4P, 3-[4-[2-(Phenoxazin-10-yl)ethoxy]phenyl]-2-butoxypropanoic
acid 267228-44-4P, Methyl
(S)-3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-methoxypropanoate
267228-45-5P, (S)-3-[4-[2-(Phenoxazin-10-yl)ethoxy]phenyl]-2-
methoxypropanoic acid 493014-71-4P,
(S)-3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-propoxypropanoic acid
493014-73-6P, (S)-3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-
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                         493014-75-8P,
(S)-3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-tert-butoxypropanoic acid
493014-77-0P, Methyl 3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-2,3-
                 493014-78-1P, Propyl
epoxypropanoate
3-[4-[2-(phenoxazin-10-y1)] ethoxy]pheny1]-2,3-epoxypropanoate
493014-79-2P, Isopropyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2,3-
epoxypropanoate
                 493014-80-5P, Butyl
3-[4-[2-(phenoxazin-10-y1)] ethoxy]phenyl]-2,3-epoxypropanoate
493014-81-6P, tert-Butyl 3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-2,3-
epoxypropanoate 493014-82-7P, Methyl
3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-hydroxypropanoate
493014-83-8P, Propyl 3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-2-
hydroxypropanoate 493014-84-9P, Isopropyl
3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-hydroxypropanoate
493014-85-0P, Butyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2-
hydroxypropanoate 493014-86-1P, tert-Butyl
3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-hydroxypropanoate
493014-37-2P, Propyl (S)-3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-
2-propoxypropanoate 493014-88-3P, Isopropyl
(S)-3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-isopropoxypropanoate
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butoxypropanoate 493014-90-7P, tert-Butyl
(S)-3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-tert-butoxypropanoate
493014-91-89, Methyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2-
methoxypropanoate 493014-92-9P, Propyl
3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-propoxypropanoate
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isopropoxypropanoate 493014-94-1P, Butyl
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ΤТ

ΤT

3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-butoxypropanoate 493014-95-2P, tert-Butyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2-tert-butoxypropanoate 493014-96-3P, 3-[4-[2-(Phenoxazin-10-y1)] ethoxy]phenyl]-2-methoxypropanoic acid 493014-97-4P, 3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-propoxypropanoicacid 493014-98-5P, 3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2isopropoxypropanoic acid 493014-99-6P, 3-[4-[2-(Phenoxazin-10-yl)ethoxy]phenyl]-2-tert-butoxypropanoic acid 493031-11-1P, (S) -3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2butoxypropanoic acid RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (intermediate; improved preparation of antidiabetic phenoxazine derivs., e.g., ragaglitazar arginine salt)

L8 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1995:229636 CAPLUS Full-text

DOCUMENT NUMBER: 123:111772 ORIGINAL REFERENCE NO.: 123:19965a

TITLE: 2,3-Diaza-1,3-dienes (Azines) as Substrates for the

Staudinger Reaction. Synthesis and Reactivity of

 $N-Imino-\beta-lactams$

AUTHOR(S): Alcaide, Benito; Miranda, Miguel; Perez-Castells,

Javier; Polanco, Concepcion; Sierra, Miguel A.

CORPORATE SOURCE: Facultad de Quimica, Universidad Complutense, Madrid,

28040, Spain

SOURCE: Journal of Organic Chemistry (1994), 59(26),

8003-10

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 123:111772

GΙ

/ Structure 101 in file .gra /

- The reaction of aromatic and aliphatic azines with different ketene AB precursors, such as the acid chloride/Et3N system, alkoxychromium(0) carbenes, and free di-Ph ketene, gives N-imino- β -lactams, e.g., I (R1 = PhO, MeO, PhCH2O, phthalimidyl, MeCO2, Me3CO, R2 = 4-MeOC6H4, Ph, 2-furyl, etc., R3 = H, Me), in good to excellent yields, with good levels of cis, trans-selectivity. A wide variety of sym.-substituted azines derived from aldehydes and ketones are compatible with the Staudinger reaction. Chiral N-imino- β -lactams derived from sym. or unsym. (mixed) chiral azines are also obtained in good yields as essentially single enantiomers (de > 95%). Different reaction intermediates, including hemiaminals, oxadiazols, and hydrazides have been isolated. Free di-Ph ketene forms Diels-Alder adducts and N-acylazadienes in addition to the previously reported N-imino- β -lactams. The usual reactivity of the β -lactam ring is modified in N-imino- β -lactams by the presence of the imino group. Thus, β -hydrazono esters, N-alkylamino- β -lactams, and NH- β -lactams can be efficiently obtained by base-catalyzed 2-azetidinone ring opening, catalytic hydrogenation, and ozonolysis, resp.
- TI 2,3-Diaza-1,3-dienes (Azines) as Substrates for the Staudinger Reaction. Synthesis and Reactivity of N-Imino- β -lactams
- SO Journal of Organic Chemistry (1994), 59(26), 8003-10

CODEN: JOCEAH; ISSN: 0022-3263

AΒ The reaction of aromatic and aliphatic azines with different ketene precursors, such as the acid chloride/Et3N system, alkoxychromium(0) carbenes, and free di-Ph ketene, gives N-imino- β -lactams, e.g., I (R1 = PhO, MeO, PhCH2O, phthalimidyl, MeCO2, Me3CO, R2 = 4-MeOC6H4, Ph, 2-furyl, etc., R3 = H, Me), in good to excellent yields, with good levels of cis, trans-selectivity. A wide variety of sym.-substituted azines derived from aldehydes and ketones are compatible with the Staudinger reaction. Chiral N-imino- β -lactams derived from sym. or unsym. (mixed) chiral azines are also obtained in good yields as essentially single enantiomers (de > 95%). Different reaction intermediates, including hemiaminals, oxadiazols, and hydrazides have been isolated. Free di-Ph ketene forms Diels-Alder adducts and N-acylazadienes in addition to the previously reported N-imino- β -lactams. The usual reactivity of the β -lactam ring is modified in N-imino- β -lactams by the presence of the imino group. Thus, β -hydrazono esters, N-alkylamino- β -lactams, and NH- β -lactams can be efficiently obtained by base-catalyzed 2-azetidinone ring opening, catalytic hydrogenation, and ozonolysis, resp.

IT 145654-57-5P 165374-81-2P 165374-82-3P 165374-83-4P 165374-90-3F 165374-91-4P 165374-92-5P

165374-93-6F 165374-94-7P 165374-95-8P 165374-96-9P

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RL: SPN (Synthetic preparation); PREP (Preparation)

(Staudinger reaction of azines and preparation and reactions of N-imino- $\beta\text{-lactams})$

L8 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1993:539588 CAPLUS Full-text

DOCUMENT NUMBER: 119:139588

ORIGINAL REFERENCE NO.: 119:25059a,25062a

TITLE: New ligands for asymmetric dihydroxylation: multiple

cinchona alkaloid units attached to a central

heterocyclic core

INVENTOR(S): Hartung, Jens; Sharpless, K. Barry

PATENT ASSIGNEE(S): Massachusetts Institute of Technology, USA

SOURCE: PCT Int. Appl., 121 pp.

.....

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

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PRIORI	TY APPLN. INFO.:			US 1991-775683	A 19911010 <
				US 1988-142692	B2 19880111 <
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W 19921006 <--

/ Structure 102 in file .gra /

Osmium-catalyzed methods of addition to an olefin are studied. In the method AΒ of asym. dihydroxylation of the present invention, an olefin, a chiral ligand, an organic solvent, an aqueous solution, a base, a ferricyanide salt and an osmium-containing compound are combined. The chiral ligand is an alkaloid or alkaloid derivative linked to an organic substituent of at least 300 daltons mol. weight through a planar aromatic spacer group. The organic substituent can be another alkaloid or alkaloid derivative With the described chiral ligands, asym. dihydroxylation of olefins with high yields and enantiomeric excesses are achieved. Thus, dihydroquinidine was treated with 1,4dichlorophthalazine in DMF containing NaH to give 1,4-bis(9'-Odihydroquinidyl)phthalazine (I). To a well stirred solution of I, a solution of potassium ferricyanide, K2CO3, OsO4 in toluene was added a Me3COH-H2O solution of 1-decene, the solution was stirred for 24 h at 0° followed by addition of sodium sulfite to give 83% 1,2-decanediol with 84% ee.

New ligands for asymmetric dihydroxylation: multiple cinchona alkaloid TIunits attached to a central heterocyclic core

WO 9307142 A1 19930415 PΙ

		TENT NO				KIND	1	DATE	3		APE	LIC	CAT	ION	NO.		D.	ATE		
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	US	526046				A		,	•	,		•				- /			010	<
	ΕP	60830	7			A1														
	ΕP	60830	7																	
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	JP	398282	29			B2		2007	0926											
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	ES	221598	39			ТЗ		2004	1016		ES	199	2-9	9214	93		1	9921	006	<
PRAI	US	1991-	7756	683		A		1991	1010	<-										
	US	1988-1	1426	592		В2		1988	0111	<-										
	US	1988-1	1590	064		A2		1988	0223	<-	-									
	US	1988-2	2500	378		A2		1988	0928	<-	-									
		1990-5							0423											
	_	1991–1		-					0423											
		1991-6							.0513											
		1992-0				W			21006											
7/ D	\cap	millm-c	· a + a	1,7700	l mc	+ had	~ ^	f add	ditio	n t	O 3	ກຸດ່	$1 \land f$	in :	220	etudi	$\sim d$	Tn	+ho	motho

Osmium-catalyzed methods of addition to an olefin are studied. In the method AB of asym. dihydroxylation of the present invention, an olefin, a chiral ligand, an organic solvent, an aqueous solution, a base, a ferricyanide salt and an osmium-containing compound are combined. The chiral ligand is an alkaloid or alkaloid derivative linked to an organic substituent of at least 300 daltons mol. weight through a planar aromatic spacer group. The organic substituent can be another alkaloid or alkaloid derivative With the described chiral

```
ligands, asym. dihydroxylation of olefins with high yields and enantiomeric
     excesses are achieved. Thus, dihydroquinidine was treated with 1,4-
     dichlorophthalazine in DMF containing NaH to give 1,4-bis(9'-O-
     dihydroquinidyl)phthalazine (I). To a well stirred solution of I, a solution
     of potassium ferricyanide, K2CO3, OsO4 in toluene was added a Me3COH-H2O
     solution of 1-decene, the solution was stirred for 24 h at 0° followed by
     addition of sodium sulfite to give 83% 1,2-decanediol with 84% ee.
    100-13-0, p-Nitrostyrene 110-57-6 300-57-2, Allylbenzene
ΤТ
               611-15-4, o-Methylstyrene
    558-37-2
                                           623-70-1
                                                     637-69-4,
                     674-76-0
                                  692-70-6
    p-Methoxystyrene
                                             695-12-5
                                                       762-63-0
                                                                  768-49-0
    932-66-1, 1-Acetylcyclohexene
                                    935-00-2
                                              1193-18-6
                                                          1746-13-0, Allyl
                  1754-62-7 2039-89-6
                                         2039-90-9
                                                      2157-18-8
    phenyl ether
                                                                  2738-19-4
    3054-95-3
               4192-77-2
                           5820-22-4
                                      7367-82-0
                                                   7642-04-8
                                                               13389-42-9
    14663-11-7
                 18448-47-0, Methyl 1-cyclohexene carboxylate
                                                               20710-38-7
                            22946-43-6 27829-72-7
    21040-45-9
                 21087-29-6
                                                       31552-04-2
    34352-92-6
                 50555-04-9
                              63511-93-3
                                          63511-95-5
                                                       66323-99-7
    67364-02-7
                 71338-71-1
                              72551-28-1
                                          78277-23-3 81703-93-7
    105018-99-3 125187-81-7 125187-82-8
                                             125187-83-9 125206-15-7
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (asym. dihydroxylation of, chiral dihydroquinidine derivative for
       osmium catalyzed)
    98-83-9, reactions
                         100-42-5, reactions
                                              103-30-0, trans-Stilbene
ΙT
    766-90-5
              771-98-2, 1-Phenylcyclohexene
                                              827-54-3, 2-Vinylnaphthalene
    872-05-9, 1-Decene
                        873-66-5 1463-04-3
                                              3901-07-3, trans-Methyl
                                    7433-56-9, trans-5-Decene
     4-methoxycinnamate
                         6714-96-1
                                                                13269-52-8,
                    13633-26-6
                                17343-88-3
                                              61142-41-4, Vinylcyclooctane
    trans-3-Hexene
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (asym. dihydroxylation of, chiral dihydroquinidine ligand for
       osmium catalyzed)
ΙΤ
    56-54-2, Quinidine
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (hydrogenation of)
ΙT
    2325-10-2P
                 16355-00-3P
                               25779-13-9P
                                             31612-63-2P
                                                          32345-64-5P
    34281-90-8P
                  35638-92-7P
                               40421-51-0P
                                             40560-98-3P
                                                           49801-14-1P
    52340-78-0P
                  53448-10-5P
                                57495-92-8P
                                             77977-74-3P
                                                           78805-31-9P
    79299-22-2P 83603-02-5P
                              84276-14-2P
                                             84518-30-9P
                                                          87827-60-9P
    87858-06-8P 88196-06-9P 89063-88-7P
                                             98464-24-5P 99881-77-3P
    108666-29-1P 108741-12-4P
                                  110549-79-6P
                                                113162-03-1P
                                                               113162-04-2P
                                  121564-12-3P 122517-80-0P
    113162-11-1P
                   115889-27-5P
    124649-67-8P
                   125132-75-4P
                                  130876-03-8P
                                                130932-13-7P
                                                               130932-14-8P
    132486-47-6P
                  135042-86-3P
                                  135042-87-4P
                                                135096-77-4P
                                                               136031-93-1P
                                                               139093-44-0P
    138890-41-2P 139093-40-6P
                                  139093-41-7P
                                                139093-42-8P
    139093-45-1P 139093-46-2P
                                  139093-47-3P
                                                139165-53-0P
                                                               139165-54-1P
    139165-55-2P 139165-56-3P
                                 139165-57-4P 139165-58-5P
                                                              139165-59-6P
    139165-60-9P 139165-61-0P
                                  139165-62-1P
                                                139165-63-2P
                                                               139165-64-3P
    139165-65-4P
                  139165-66-5P
                                                139165-68-7P
                                                               139165-69-8P
                                  139165-67-6P
    139165-70-1P
                   143536-11-2P
                                  149519-66-4P
                                                149519-67-5P 149519-68-6P
    149519-69-7P
                 149562-10-7P
                                  149562-11-8P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
Г8
    ANSWER 5 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                        1992:486257 CAPLUS Full-text
DOCUMENT NUMBER:
                        117:86257
ORIGINAL REFERENCE NO.: 117:14967a,14970a
TITLE:
                        A lipid-lipase aggregate with ether linkage as a new
                        type of immobilized enzyme for enantioselective
                        hydrolysis in organic solvents
                        Akita, Hiroyuki; Umezawa, Isao; Matsukura, Hiroko;
AUTHOR(S):
```

Oishi, Takeshi

CORPORATE SOURCE: Sch. Pharm. Sci., Toho Univ., Funabashi, 274, Japan

SOURCE: Chemical & Pharmaceutical Bulletin (1992),

40(2), 318-24

CODEN: CPBTAL; ISSN: 0009-2363

DOCUMENT TYPE: Journal LANGUAGE: English

For the purpose of carrying out smoothly enzymic reaction of water-insol. substrates in organic solvents, a new type of immobilized enzyme, a lipid-lipase aggregate, was developed. To prepare various kinds of lipid-lipase aggregates, 27 kinds of dialkyl ether-type phospholipid analogs were newly synthesized and used for the preparation of aggregates with lipase. Thus obtained lipid-lipase aggregates catalyzed the enantioselective hydrolysis of the $(\pm)-\alpha$ -acyloxy ester 2 much more efficiently than lipase immobilized with synthetic prepolymer (ENTP-4000) in water-saturated iso-Pr ether. The reaction time became much shorter (2 to 3 days for completion as compared with 21 days) and the chemical and optical yields of the reaction products were high.

TI A lipid-lipase aggregate with ether linkage as a new type of immobilized enzyme for enantioselective hydrolysis in organic solvents

SO Chemical & Pharmaceutical Bulletin (1992), 40(2), 318-24 CODEN: CPBTAL; ISSN: 0009-2363

AB For the purpose of carrying out smoothly enzymic reaction of water-insol. substrates in organic solvents, a new type of immobilized enzyme, a lipid-lipase aggregate, was developed. To prepare various kinds of lipid-lipase aggregates, 27 kinds of dialkyl ether-type phospholipid analogs were newly synthesized and used for the preparation of aggregates with lipase. Thus obtained lipid-lipase aggregates catalyzed the enantioselective hydrolysis of the $(\pm)-\alpha$ -acyloxy ester 2 much more efficiently than lipase immobilized with synthetic prepolymer (ENTP-4000) in water-saturated iso-Pr ether. The reaction time became much shorter (2 to 3 days for completion as compared with 21 days) and the chemical and optical yields of the reaction products were high.

IT 64599-78-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and hydrogenation of)

IT 125354-94-1P 125354-96-3P

RL: PREP (Preparation)

(preparation of, by lipid-lipase aggregate hydrolysis in solvents)

L8 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1990:406196 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 113:6196

ORIGINAL REFERENCE NO.: 113:1203a,1206a

TITLE: Synthesis of threo-3-aryl-2,3-dihydroxypropanoic acid

derivatives with high optical purity

AUTHOR(S): Matthews, Barry R.; Gountzos, Helen; Jackson, W. Roy;

Watson, Keith G.

CORPORATE SOURCE: Dep. Chem., Monash Univ., Clayton, 3168, Australia

SOURCE: Tetrahedron Letters (1989), 30(38), 5157-8

CODEN: TELEAY; ISSN: 0040-4039

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 113:6196

GΙ

- Cyanohydrins of aromatic aldehydes can be obtained in high optical purity using the Inoue dipeptide catalyst system and converted into enantiomerically and diastereochem. pure threo-3-aryl-2,3-dihydroxypropanoic acid derivs. via a route which involves a base-catalyzed equilibration of the acetonides of the cyano diols. Thus, (R)-(+)-4-MeOC6H4CH(OH)CN was hydrogenated over Ni and treated sequentially with NaHSO3, NaCN, and (MeO)2CMe2 and acid to give cisand trans-dioxolanes I. Hydrolysis and epimerization of I with KOH in EtOH and then treatment with aqueous acid followed by HCl-MeOH gave the enantiomerically pure threo-4-MeOC6H4CH(OH)CH(OH)CO2Me (II). II was successfully converted into diltiazem.
- TI Synthesis of threo-3-aryl-2,3-dihydroxypropanoic acid derivatives with high optical purity
- TI Synthesis of threo-3-aryl-2,3-dihydroxypropanoic acid derivatives with high optical purity
- SO Tetrahedron Letters (1989), 30(38), 5157-8 CODEN: TELEAY; ISSN: 0040-4039
- Cyanohydrins of aromatic aldehydes can be obtained in high optical purity using the Inoue dipeptide catalyst system and converted into enantiomerically and diastereochem. pure threo-3-aryl-2,3-dihydroxypropanoic acid derivs. via a route which involves a base-catalyzed equilibration of the acetonides of the cyano diols. Thus, (R)-(+)-4-MeOC6H4CH(OH)CN was bydrogenated over Ni and treated sequentially with NaHSO3, NaCN, and (MeO)2CMe2 and acid to give cisand trans-dioxolanes I. Hydrolysis and epimerization of I with KOH in EtOH and then treatment with aqueous acid followed by HCl-MeOH gave the enantiomerically pure threo-4-MeOC6H4CH(OH)CH(OH)CO2Me (II). II was successfully converted into diltiazem.
- IT 97070-73-0

RL: RCT (Reactant); RACT (Reactant or reagent)
 (hydrogenation and cyanation of)

IT 122517-80-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and conversion of, to diltiazem)

L8 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1990:75971 CAPLUS Full-text

DOCUMENT NUMBER: 112:75971

ORIGINAL REFERENCE NO.: 112:12975a,12978a

TITLE: Ligand-accelerated catalytic asymmetric

dihydroxylation

INVENTOR(S): Marko, Istvan E.; Sharpless, K. Barry

PATENT ASSIGNEE(S): Massachusetts Institute of Technology, USA

SOURCE: PCT Int. Appl., 66 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 8906225 W: JP	A1 19890713	WO 1989-US86	19890110 <
RW: AT, BE, CH,	DE, FR, GB, IT, LU	, NL, SE	
US 4871855	A 19891003	US 1988-159068	19880223 <
US 4965364	A 19901023	US 1988-250378	19880928 <
EP 39 572 9	A1 19901107	EP 1989-901900	19890110 <
EP 39 572 9	B1 19950927		
R: AT, BE, CH,	DE, FR, GB, IT, LI	, LU, NL, SE	
JP 03503885	T 19910829	JP 1989-501814	19890110 <
JP 3153540	B2 20010409		

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EP 658532
                        A1
                                19950621 EP 1995-200458
                                                                   19890110 <--
     EP 658532
                         В1
                               19990407
        R: AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE
                                                                   19890110 <--
     AT 128449
                     T 19951015 AT 1989-901900
     AT 178578
                         Τ
                                19990415
                                            AT 1995-200458
                                                                   19890110 <--
     JP 2001192383
                         Α
                                20010717
                                           JP 2000-335775
                                                                   19890110 <--
     CA 1338314
                        С
                                19960507
                                            CA 1989-587964
                                                                   19890111 <--
                                            US 1988-142692 A 19880111 <--

US 1988-159068 A 19880223 <--

US 1988-250378 A 19880928 <--
PRIORITY APPLN. INFO.:
                                            EP 1989-901900
                                                               A3 19890110 <--
                                            JP 1989-501814
                                                                A3 19890110 <--
                                            WO 1989-US86
                                                                W 19890110 <--
```

Vicinal diols are prepared by asym. dihydroxylation of olefins in the presence of a chiral ligand, an Os-containing catalyst, an amine oxide, an organic solvent, and H2O. OsO4 was injected into a solution of trans-stilbene, hydroquinidine p-chlorobenzoate, and N-methylmorpholine N-oxide in Me2CO and H2O at $0-4^{\circ}$ with shaking, Na2S2O5 added, and the mixture worked up to give 55% (R,R)-PhCH(OH)CH(OH)Ph of 99% enantiomeric excess. Preparation of dihydroquinidine derivs. and their recovery were also given. Asym. dihydroxylations of trans-3-hexene, 1-phenylcyclohexene, β -methylstyrene, and Me (E)-4-methoxycinnamate were also given.

TI Ligand-accelerated catalytic asymmetric dihydroxylation

PΙ	WO	8906225	Α1	19890713

LI	PAT	ENT NO.		KIND		APPLICATION NO.	
PI	WO	8906225 W: JP				WO 1989-US86	19890110 <
		RW: AT, BE,	CH,	DE, FI	R, GB, IT,	LU, NL, SE	
						US 1988-159068	19880223 <
	US	4965364		A	19901023	US 1988-250378	19880928 <
	ΕP	395729		A1	19901107	EP 1989-901900	19890110 <
	ΕP	395729		B1	19950927		
						LI, LU, NL, SE	
						JP 1989-501814	19890110 <
	JΡ	3153540		В2	20010409		
						EP 1995-200458	19890110 <
		658532					
				•		LI, LU, NL, SE	
						AT 1989-901900	
	ΑT	178578		T	19990415	AT 1995-200458	19890110 <
	JΡ	2001192383		A	20010717	JP 2000-335775	19890110 <
	CA	1338314		С	19960507	CA 1989-587964	19890111 <
PRAI		1988-142692				<	
	US	1988-159068		А	19880223	<	
	US	1988-250378		A	19880928	<	
	EΡ	1989-901900		A3	19890110	<	
		1989-501814			19890110	<	
	WO	1989-US86		W	19890110	<	

Vicinal diols are prepared by asym. dihydroxylation of olefins in the presence of a chiral ligand, an Os-containing catalyst, an amine oxide, an organic solvent, and H2O. OsO4 was injected into a solution of trans-stilbene, hydroquinidine p-chlorobenzoate, and N-methylmorpholine N-oxide in Me2CO and H2O at $0-4^{\circ}$ with shaking, Na2S2O5 added, and the mixture worked up to give 55% (R,R)-PhCH(OH)CH(OH)Ph of 99% enantiomeric excess. Preparation of dihydroquinidine derivs. and their recovery were also given. Asym. dihydroxylations of trans-3-hexene, 1-phenylcyclohexene, β -methylstyrene, and Me (E)-4-methoxycinnamate were also given.

ST asym dihydroxylation olefin chiral ligand

IT Hydroxylation

(osmylation, stereoselective, of olefins, chiral ligands as accelerators for)

IT 56-54-2

RL: RCT (Reactant); RACT (Reactant or reagent)

(hydrogenation of)

IT 40421-51-0P 52340-78-0P 53448-10-5P 125073-64-5P

125132-75-4P

L8 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1987:534332 CAPLUS Full-text

DOCUMENT NUMBER: 107:134332

ORIGINAL REFERENCE NO.: 107:21708h,21709a

TITLE: Preparation of 1,5-benzothiazepin-4-one derivatives as

blood platelet aggregation inhibitors

INVENTOR(S): Inoue, Hirozumi; Otsuka, Hisao PATENT ASSIGNEE(S): Tanabe Seiyaku Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 15 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 62096482	A	19870502	JP 1986-141592	19860617 <
JP 05081597	В	19931115		
PRIORITY APPLN. INFO.:			JP 1985-136569	A1 19850621 <
GT				

/ Structure 104 in file .gra /

- The title compds. [I, II and III; R1 = alkyl, alkoxy; R2 = H, alkanoyl; R3 = alkyl; R4 = H; R5 or R6 is H and the other is C1; R7 or R8 is alkyl, alkoxy, alkylthio, F, PhCO2O or OH and the other is H, or R7 = R8 = alkoxy], useful as blood platelet aggregation inhibitors, were prepared by treatment of I, II and III (R4 = alkyl) with acid halides of haloformic acids, e.g., COCl2, or their esters, e.g., CCl3O2CCl, and deacylation of the resulting I, II and III (R4 = COR9; R9 = halo or ester residue). COCl2 (31.91 g) in toluene was added to a solution of 20.21 g II (R1 = OMe, R2 = Ac, R3 = R4 = Me, R5 = C1, R6 = H) in toluene and the mixture was stirred for 16 h at 30° to give II (R1 = OMe, R2 = Ac, R3 = Me, R4 = COCl, R5 = Cl, R6 = H), which was treated with 10% aqueous HCl and MeCN under reflux to give II (R1 = OMe, R2 = R4 = R6 = H, R3 = Me, R5 = Cl) (as the fumarate). In vitro, this inhibited collagen-induced blood platelet aggregation.
- TI Preparation of 1,5-benzothiazepin-4-one derivatives as blood platelet aggregation inhibitors

	aggregacion im	TOTCOLD			
ΡI	JP 62096482 A	19870502	Showa		
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PΙ	JP 620964 8 2	A	19870502	JP 1986-141592	19860617 <
	JP 05081597	В	19931115		
PRAI	JP 19 85 -136569	A1	19850621	<	
ΙT	103921-06-8P				

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RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and cyclization of, dihydrobenzothiazepine derivative from)
ΤТ
     103921-05-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and hydrogenation of)
     100902-58-7P, (±)-cis-2-(4-Methoxyphenyl)-3-hydroxy-9-chloro-2,3-
ΤТ
     dihydro-1,5-benzothiazepine-4(5H)-one
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and resolution of, via chiral
        (naphthylsulfonyl)pyrrolidinecarboxylaye ester)
=> S L6 AND (Ru? or Rh? or Pd? or Os?)
      1446878 RU?
       783851 RH?
       329541 PD?
       581552 OS?
           18 L6 AND (RU? OR RH? OR PD? OR OS?)
L9
=> d 19 1-5 ibib abs ti hit
    ANSWER 1 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2007:544712 CAPLUS <u>Full-text</u>
DOCUMENT NUMBER:
                        148:449648
TITLE:
                        An improved process for the preparation of phenoxazine
                        antidiabetic compounds
INVENTOR(S):
                        Rao, Siripragada Mahender; Reddy, Chepyala Naveen
                        Kumar; Reddy, Challa Maheedhara; Sarma, Mamillapalli
                        Ramabhandra; Reddy, Gaddam Om
PATENT ASSIGNEE(S):
                        Reddy's Laboratories Ltd., India
SOURCE:
                        Indian Pat. Appl., 18pp.
                        CODEN: INXXBQ
DOCUMENT TYPE:
                        Patent
                        English
LANGUAGE:
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
                                         APPLICATION NO.
     PATENT NO.
                       KIND
                               DATE
                       ____
                                          _____
                                                                 _____
    IN 2002MA00508
                                         IN 2002-MA508
IN 2002-MA508
                       A 20070511
                                                                20020708 <--
PRIORITY APPLN. INFO.:
                                                                20020708 <--
OTHER SOURCE(S): CASREACT 148:449648; MARPAT 148:449648
GI
```

/ Structure 105 in file .gra /

The invention relates to an improved process for the preparation of antidiabetic phenoxazinylethoxyphenylalkoxypropanoate arginine salts I [R1 = Me, Et, Pr, i-Pr]. The process uses inexpensive chems. and an easy resolution via chiral amine salts. Thus, condensation of 4-[2-(phenoxazin-10-yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the corresponding glycidic ester, which was hydrogenated over Pd/C, O-ethylated with di-Et sulfate in xylene, and saponified with aqueous NaOH in MeOH to give

acid (\pm) -II. Resolution of this racemate using (-)-ephedrine and salification with L-arginine in i-PrOH gave I (R1 = Et).

TΙ An improved process for the preparation of phenoxazine antidiabetic

compounds

APPLICATION NO. KIND DATE PATENT NO. _____ ____ -----_____ _____ IN 2002MA00508 PΙ A 20070511 IN 2002-MA508 20020708 <--PRAI IN 2002-MA508 20020708 <--

The invention relates to an improved process for the preparation of antidiabetic phenoxazinylethoxyphenylalkoxypropanoate arginine salts I [R1 = Me, Et, Pr, i-Pr]. The process uses inexpensive chems. and an easy resolution via chiral amine salts. Thus, condensation of 4-[2-(phenoxazin-10yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the corresponding glycidic ester, which was hydrogenated over Fd/C, O-ethylated with di-Et sulfate in xylene, and saponified with aqueous NaOH in MeOH to give acid (±)-II. Resolution of this racemate using (-)-ephedrine and salification with L-arginine in i-PrOH gave I (R1 = Et).

222834-12-0P 222834-14-2P 222834-21-1P 222834-30-2P ΤT 493014-69-0P 493014-71-4P 493014-73-6P 493014-77-0P 267228-45-5P 493014-78-1P 493014-79-2P 493014-82-7P 493014-83-8P 493014-84-9P 493014-96-3P 493014-97-4P 493014-98-5P RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation); RACT (Reactant or reagent)

(improved preparation of phenoxazine antidiabetic compds.)

ANSWER 2 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:242286 CAPLUS Full-text

DOCUMENT NUMBER: 138:271396

Preparation of 3-aryl-2-alkoxypropanoates from TITLE:

3-aryl-2-oxopropanoates via ketalization and reduction

INVENTOR(S): Siripragada, Mahender Rao; Vanadanapu, Loka Appala

Purushotham; Mamillapalli, Ramabhadra Sarma; Gaddam,

Om Reddy

PATENT ASSIGNEE(S): Reddy's Laboratories Ltd., India

SOURCE: PCT Int. Appl., 12 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	PATENT NO.						KIND DATE			APPLICATION NO.									
WO	2003	0249	 15		A1 20030327								20020919 <						
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,		
		СО,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,		
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,		
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,		
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THER S	HER SOURCE(S).					CASPEACT 138.271396. MARPAT 138.271396													

OTHER SOURCE(S): CASREACT 138:271396; MARPAT 138:271396 AB 4-HOC6H4CH2CH(OR1)CO2R2 (R1 = H, alkyl; R2 = alkyl), were prepared by converting 4-HOC6H4CH2COCO2H to 4-HOC6H4CH2C(OR1)2CO2R2 (variables as above) in the presence of acid followed by reduction in HOAc at 40-80 psi for 6-12 h. Thus, 4-hydroxyphenylpyruvic acid was stirred in EtOH at 10-15° for 3 h, at room temperature for 6 h, and at 50-60° for 4 h to give 4-HOC6H4CH2C(OEt)2CO2Et. The latter was hydrogenated in EtOH/HOAc over Rh/Al2O3 at 60 psi for 12 h to give 4-HOC6H4CH2CH(OEt)CO2Et.

TI Preparation of 3-aryl-2-alkoxypropanoates from 3-aryl-2-oxopropanoates via ketalization and reduction

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REFERENCE COUNT:
                             THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
                             RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
    PATENT NO.
                       KIND
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                                         APPLICATION NO.
                              _____
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                              20030327 WO 2002-IB3874
    WO 2003024915
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PI
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AB 4-HOC6H4CH2CH(OR1)CO2R2 (R1 = H, alkyl; R2 = alkyl), were prepared by converting 4-HOC6H4CH2COCO2H to 4-HOC6H4CH2C(OR1)2CO2R2 (variables as above) in the presence of acid followed by reduction in HOAc at 40-80 psi for 6-12 h. Thus, 4-hydroxyphenylpyruvic acid was stirred in EtOH at 10-15° for 3 h, at room temperature for 6 h, and at 50-60° for 4 h to give 4-HOC6H4CH2C(OEt)2CO2Et. The latter was hydrogenated in EtOH/HOAc over Rh/Al2O3 at 60 psi for 12 h to give 4-HOC6H4CH2CH(OEt)CO2Et.

IT 1314-15-4, Platinum dioxide 7440-02-0, Nickel, uses 7440-16-6, Rhodium, uses

RL: CAT (Catalyst use); USES (Uses)

(preparation of 3-aryl-2-alkoxypropanoates from <math>3-aryl-2-oxopropanoates via ketalization and reduction)

IT 4375-92-2P

RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of 3-aryl-2-alkoxypropanoates from 3-aryl-2-oxopropanoates via ketalization and reduction)

IT 197299-16-4P

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREF (Preparation)

(preparation of 3-aryl-2-alkoxypropanoates from <math>3-aryl-2-oxopropanoates via ketalization and reduction)

L9 ANSWER 3 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:76765 CAPLUS Full-text

DOCUMENT NUMBER: 138:137318

TITLE: An improved process for the preparation of

antidiabetic phenoxazine compounds, e.g., ragaglitazar

L-arginine salt

INVENTOR(S): Siripragada, Mahender Rao; Chepyala, Naveen Kumar

Reddy; Challa, Maheedharareddy; Mamillapalli,

Ramabhadra Sarma; Gaddam, Om Reddy

PATENT ASSIGNEE(S): Reddy's Laboratories Ltd., India

SOURCE: PCT Int. Appl., 31 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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M(WO 2003008397						 2003	0130	WO 2002-IB2776						20020716 <				
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OTHER :	OTHER SOURCE(S):					REAC	T 13	8:13	7318; MARPAT 138:137318					318					

/ Structure 106 in file .gra /

- The invention relates to an improved, multi-step process for the preparation AΒ of antidiabetic compds. I [where R1 represents an alkyl group such as Me, Et, Pr, iso-Pr, Bu, tert-Bu, and the like]. I are known antidiabetic, hypolipidemic, and antihypercholesterolemic agents. Older methods of preparing I are industrially unsuitable for a variety of reasons, including exotic and moisture-sensitive reactions, tedious purifns., low yields, and long cycle times. The new method is simple, robust, scalable, and provides I with high chemical and chiral purity. The use of highly reactive, difficultto-handle, and expensive chems. is avoided, and simple, inexpensive chems. such as di-Et sulfate and K2CO3 are used instead. Two variants of the process are claimed and illustrated, in which the resolution step occurs at different points. For instance, Darzen's condensation of 4-[2-(phenoxazin-10yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the corresponding glycidic ester, which was hydrogenated over Pd/C and saponified with aqueous NaOH in MeOH to give acid (±)-II. Two-step resolution of this racemate using (S)-phenylglycinol and then (-)- α -methylbenzylamine gave (S)-II of 99.2% purity. Combined etherification/esterification of the latter with NaH and EtI, saponification of the ester with aqueous NaOH in MeOH, and salification with L-arginine in iso-PrOH gave I (R1 = Et), i.e., the arginine salt of ragaglitazar.. The alternative sequence involves Darzen's condensation, hydrogenation, etherification/esterification, ester saponification, resolution, and salification.
- TI An improved process for the preparation of antidiabetic phenoxazine compounds, e.g., ragaglitazar L-arginine salt
 REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS

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RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
    PATENT NO.
                       KIND
                              DATE APPLICATION NO. DATE
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                               20030130 WO 2002-IB2776
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                        A1
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            PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
            UA, UG, US, UZ, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
            CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
            PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
            NE, SN, TD, TG
                               20031220 IN 2001-MA585
    IN 191728
                        A1
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    AU 2002313569
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PRAI IN 2001-MA585
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    WO 2002-IB2776
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                               20020716 <--
     The invention relates to an improved, multi-step process for the preparation
AΒ
     of antidiabetic compds. I [where R1 represents an alkyl group such as Me, Et,
     Pr, iso-Pr, Bu, tert-Bu, and the like]. I are known antidiabetic,
     hypolipidemic, and antihypercholesterolemic agents. Older methods of
     preparing I are industrially unsuitable for a variety of reasons, including
     exotic and moisture-sensitive reactions, tedious purifns., low yields, and
     long cycle times. The new method is simple, robust, scalable, and provides I
     with high chemical and chiral purity. The use of highly reactive, difficult-
     to-handle, and expensive chems. is avoided, and simple, inexpensive chems.
     such as di-Et sulfate and K2CO3 are used instead. Two variants of the process
     are claimed and illustrated, in which the resolution step occurs at different
     points. For instance, Darzen's condensation of 4-[2-(phenoxazin-10-
     yl)ethoxy]benzaldehyde with Et chloroacetate using EtONa in EtOH gave the
     corresponding glycidic ester, which was hydrogenated over Pd/C and saponified
     with aqueous NaOH in MeOH to give acid (\pm)-II. Two-step resolution of this
     racemate using (S)-phenylglycinol and then (-)-\alpha-methylbenzylamine gave (S)-II
     of 99.2% purity. Combined etherification/esterification of the latter with
     NaH and EtI, saponification of the ester with aqueous NaOH in MeOH, and
     salification with L-arginine in iso-PrOH gave I (R1 = Et), i.e., the arginine
     salt of ragaglitazar.. The alternative sequence involves Darzen's
     condensation, hydrogenation, etherification/esterification, ester
     saponification, resolution, and salification.
    232834-12-0P, Ethyl 3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-2-
ΙT
    ethoxypropanoate 222834-14-2P, Ethyl
    3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-2-hydroxypropanoate
    222834-21-1P, 3-[4-[2-(Phenoxazin-10-yl)ethoxy]phenyl]-2-ethoxypropanoic
    acid
           222834-23-3P, 3-[4-[2-(Phenoxazin-10-yl)ethoxy]phenyl]-2-
                           222834-30-2P,
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    RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic
    preparation); PREP (Preparation); RACT (Reactant or reagent)
        (intermediate; improved preparation of antidiabetic phenoxazine derivs.,
        e.g., ragaglitazar arginine salt)
IT
    222834-24-4P, 3-[4-[2-(Phenoxazin-10-yl)ethoxy]phenyl]-2-butoxypropanoic
    acid 267228-44-4P, Methyl
     (S)-3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-methoxypropanoate
    267228-45-5P, (S)-3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-
    methoxypropanoic acid 493014-71-4P,
     (S)-3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-propoxypropanoic acid
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isopropoxypropanoic acid
                              493014-75-8P,
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     493014-77-0P, Methyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2,3-
                     493014-78-1P, Propyl
     epoxypropanoate
     3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2,3-epoxypropanoate
     493014-79-2P, Isopropyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2,3-
                      493014-80-5P, Butyl
     epoxypropanoate
     3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2,3-epoxypropanoate
     493014-81-6P, tert-Butyl 3-[4-[2-(phenoxazin-10-v1)ethoxy]phenvl]-2,3-
     epoxypropanoate 493014-82-7P, Methyl
     3-[4-[2-(phenoxazin-10-y1)ethoxy]pheny1]-2-hydroxypropanoate
     493014-83-8P, Propyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2-
     hydroxypropanoate 493014-84-9P, Isopropyl
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     493014-85-09, Butyl 3-[4-[2-(phenoxazin-10-y1)ethoxy]phenyl]-2-
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     3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-2-hydroxypropanoate
     493014-87-2P, Propyl (S)-3-[4-[2-(phenoxazin-10-yl)ethoxy]phenyl]-
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     3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-tert-butoxypropanoic acid
     493031-11-1P, (S)-3-[4-[2-(Phenoxazin-10-y1)ethoxy]phenyl]-2-
     butoxypropanoic acid
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (intermediate; improved preparation of antidiabetic phenoxazine derivs.,
        e.g., ragaglitazar arginine salt)
L9
    ANSWER 4 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                        1996:262036 CAPLUS Full-text
DOCUMENT NUMBER:
                         124:289000
ORIGINAL REFERENCE NO.: 124:53583a,53586a
TITLE:
                        Aydrogenation process and catalysts for the
                         production of phenyllactic acids from phenylpyruvic
                         Morita, Hikari; Mori, Hiroyuki
INVENTOR(S):
PATENT ASSIGNEE(S):
                         Nitto Chemical Industry Co., Ltd., Japan
SOURCE:
                         Eur. Pat. Appl., 8 pp.
                         CODEN: EPXXDW
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
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493014-73-6P, (S)-3-[4-[2-(Phenoxazin-10-y1)ethoxy]pheny1]-2-

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EP 696566	A1	19960214	EP 1995-112540	_	19950809	<
EP 696566	В1	19980610				
R: CH, DE, LI						
JP 08053394	A	19960227	JP 1994-206102		19940809	<
JP 3394606	B2	20030407				
US 5684186	A	19971104	US 1995-511152		19950804	<
CN 1122325	A	19960515	CN 1995-109050		19950809	<
CN 1083422	С	20020424				
PRIORITY APPLN. INFO.:			JP 1994-206102	A	19940809	<
OTHER SOURCE(S):	CASRE	ACT 124:2890	00; MARPAT 124:289000			
GI						

/ Structure 107 in file .gra /

- AB Phenyllactic acids [I; R1, R2 = H, OH, (un)branched alkyl, (un)branched alkoxy; R3 = H, (un)branched alkyl; R1R2 = methylenedioxy], useful as intermediates in the preparation of agrochems. (no data) and pharmaceuticals (no data), are prepared in high yield by the hydrogenation of phenylpyruvic acids (II) in the presence of a catalyst containing ≥1 Group VIII metal (e.g., Pd, Pt, Ni, etc.). Thus, 3-(4-hydroxyphphenyl)pyruvic acid was hydrogenated in MeOH using a Pd/C catalyst at 25°/5 kg/cm2, producing 3-(4-hydroxyphenyl)lactic acid.
- TI Hydrogenation process and catalysts for the production of phenyllactic acids from phenylpyruvic acids
- TI Hydrogenation process and catalysts for the production of phenyllactic acids from phenylpyruvic acids
- PI EP 696566 A1 19960214

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ΡI	EP 696566	A1	19960214	EP 1995-112540	19950809 <
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- AB Phenyllactic acids [I; R1, R2 = H, OH, (un)branched alkyl, (un)branched alkoxy; R3 = H, (un)branched alkyl; R1R2 = methylenedioxy], useful as intermediates in the preparation of agrochems. (no data) and pharmaceuticals (no data), are prepared in high yield by the hydrogenation of phenylpyruvic acids (II) in the presence of a catalyst containing ≥1 Group VIII metal (e.g., Pd, Pt, Ni, etc.). Thus, 3-(4-hydroxyphenyl)pyruvic acid was hydrogenated in MeOH using a Pd/C catalyst at 25°/5 kg/cm2, producing 3-(4-hydroxyphenyl)lactic acid.
- ST phenyllactic acid prepn; hydroxyphenyllactic acid prepn; phenylpyruvic hydrogenation prepn phenyllactic acid
- IT Aydrogenation catalysts

(Group VIII metals for the conversion of phenylpyruvic acids into phenyllactic acids)

- IT Group VIII elements
 - RL: CAT (Catalyst use); USES (Uses)

(hydrogenation catalysts for the production of phenyllactic acids from phenylpyruvic acids)

IT Hydrogenation

(of phenylpyruvic acids in the production of phenyllactic acids) ΙT 7440-02-0, Nickel, uses 7440-05-3, Palladium, uses 7440-06-4, Platinum, uses 7440-16-6, Phodium, uses 7440-48-4, Cobalt, uses RL: CAT (Catalyst use); USES (Uses) (bydrogenation process and catalysts for the production of phenyllactic acids from phenylpyruvic acids) 123-08-0, 4-Hydroxybenzaldehyde 156-06-9 543-24-8, N-Acetylglycine 884-18-4 1201-77-0 4228-66-4 4607-41-4 28030-16-2 38335-22-7 ΤТ 39829-17-9 69882-69-5 70028-57-8 100612-75-7 115863-73-5 175897-64-0 175897-71-9 RL: RCT (Reactant); RACT (Reactant or reagent) (hydrogenation process and catalysts for the production of phenyllactic acids from phenylpyruvic acids) 156-39-8P, 3-(4-Hydroxyphenyl)pyruvic acid 38243-39-9P 52507-17-2P ΤT RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (bydrogenation process and catalysts for the production of phenyllactic acids from phenylpyruvic acids) 306-23-0P, 3-(4-Hydroxyphenyl)lactic acid 828-01-3P 949-14-4P ΙT 3247-74-3P 6803-09-4P 23028-17-3P 28030-15-1P 52262-43-8P 55301-58-1P 175897-65-1P 175897-66-2P 175897-67-3P 175897-68-4P 175897-69-5P 175897-70-8P RL: SPN (Synthetic preparation); PREP (Preparation) (hydrogenation process and catalysts for the production of phenyllactic acids from phenylpyruvic acids) ANSWER 5 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN L9ACCESSION NUMBER: 1993:539588 CAPLUS <u>Full-text</u> DOCUMENT NUMBER: 119:139588 ORIGINAL REFERENCE NO.: 119:25059a,25062a New ligands for asymmetric dihydroxylation: multiple TITLE: cinchona alkaloid units attached to a central heterocyclic core Hartung, Jens; Sharpless, K. Barry INVENTOR(S): Massachusetts Institute of Technology, USA PATENT ASSIGNEE(S): PCT Int. Appl., 121 pp. SOURCE: CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 5 PATENT INFORMATION: KIND DATE APPLICATION NO. PATENT NO. ----_____ WO 9307142 A1 19930415 WO 1992-US8544 19921006 <--W: CA, JP RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, SE US 5260461 A 19931109 US 1991-775683 19911010 <--EP 608307 A1 19940803 EP 1992-921493 19921006 <--B1 20040204 EP 608307 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, SE R: AT, BE, CH, DE, DK, ES, FK, GB, GK, IE, II, LI, LO, FC, NL, SE

JP 07500323

T 19950112

JP 1993-507170

19921006 <-
JP 3982829

CA 2120919

C 20030701

CA 1992-2120919

AT 258932

T 20040215

AT 1992-921493

19921006 <-
ES 2215989

RITY APPLN. INFO.:

US 1988-142692

US 1988-159064

A2 19880223 <--

PRIORITY APPLN. INFO.:

/ Structure 108 in file .gra /

AB Osmium-catalyzed methods of addition to an olefin are studied. In the method of asym. dihydroxylation of the present invention, an olefin, a chiral ligand, an organic solvent, an aqueous solution, a base, a ferricyanide salt and an osmium-containing compound are combined. The chiral ligand is an alkaloid or alkaloid derivative linked to an organic substituent of at least 300 daltons mol. weight through a planar aromatic spacer group. The organic substituent can be another alkaloid or alkaloid derivative With the described chiral ligands, asym. dihydroxylation of olefins with high yields and enantiomeric excesses are achieved. Thus, dihydroquinidine was treated with 1,4-dichlorophthalazine in DMF containing NaH to give 1,4-bis(9'-O-dihydroquinidyl)phthalazine (I). To a well stirred solution of I, a solution of potassium ferricyanide, K2CO3, OsO4 in toluene was added a Me3COH-H2O solution of 1-decene, the solution was stirred for 24 h at 0° followed by addition of sodium sulfite to give 83% 1,2-decanediol with 84% ee.

TI New ligands for asymmetric dihydroxylation: multiple cinchona alkaloid units attached to a central heterocyclic core

PI WO 9307142 A1 19930415

		TENT NO.															ATE		
PI		9307142 W: CA,															9921	006	<
		RW: AT,		CH,	DE,	DK,	ES,	FR,	GB,	GF	R, IE	Ξ,	IT,	LU,	MC,	NL,	SE		
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	US	1988-142	2692		В2	1	1988	0111	<-	-									
	US	1988-159	064		A2	1	1988	0223	<-	_									
	US	1988-250	378		A2	1	1988	0928	<-	-									
	US	1990-512	2934		A2	1	1990	0423	<-	_									
	WO	1991-US2	2778		W	1	1991	0423	<-	_									
	US	1991-699	183		A2	1	1991	0513	<-	_									
	WO	1992-US8	3544		\overline{W}	1	1992	1006	<-	_									
ND	100	milion cat	211770	dmo	+ hode	- of	200	14+40	n +		2 2	of.	in -		0+1101	\sim d	Tη	+ho	motho

AB Osmium-catalyzed methods of addition to an olefin are studied. In the method of asym. dihydroxylation of the present invention, an olefin, a chiral ligand, an organic solvent, an aqueous solution, a base, a ferricyanide salt and an osmium-containing compound are combined. The chiral ligand is an alkaloid or alkaloid derivative linked to an organic substituent of at least 300 daltons mol. weight through a planar aromatic spacer group. The organic substituent can be another alkaloid or alkaloid derivative With the described chiral ligands, asym. dihydroxylation of olefins with high yields and enantiomeric

excesses are achieved. Thus, dihydroquinidine was treated with 1,4dichlorophthalazine in DMF containing NaH to give 1,4-bis(9'-Odihydroquinidyl)phthalazine (I). To a well stirred solution of I, a solution of potassium ferricyanide, K2CO3, O3O4 in toluene was added a Me3COH-H2O solution of 1-decene, the solution was stirred for 24 h at 0° followed by addition of sodium sulfite to give 83% 1,2-decanediol with 84% ee. asym dihydroxylation catalyst dihydroquinidine ligand; alkene dihydrozylation catalyst; cinchona alkaloid osmylation catalyst Asymmetric synthesis and induction (osmium catalyzed dihydroxylation, cinchona alkaloid derivs. as accelerator for) Hydroxylation catalysts (osmylation, stereoselective, ligand accelerated, dihydroquinidine derivs. for) 110-57-6 300-57-2, Allylbenzene 447-53-0 100-13-0, p-Nitrostyrene 558-37-2 611-15-4, o-Methylstyrene 623-70-1 637-69-4, p-Methoxystyrene 674-76-0 692-70-6 695-12-5 762-63-0 768-49-0 932-66-1, 1-Acetylcyclohexene 935-00-2 1193-18-6 1746-13-0, Allyl phenyl ether 1754-62-7 2039-89-6 2039-90-9 2157-18-8 2738-19-4 7367-82-0 5820-22-4 7642-04-8 3054-95-3 4192-77-2 13389-42-9 20710-38-7 14663-11-7 18448-47-0, Methyl 1-cyclohexene carboxylate 21087-29-6 22946-43-6 27829-72-7 31552-04-2 21040-45-9 34352-92-6 50555-04-9 63511-93-3 63511-95-5 66323-99-7 67364-02-7 71338-71-1 72551-28-1 78277-23-3 81703-93-7 105018-99-3 125187-83-9 125187-81-7 125187-82-8 125206-15-7 RL: RCT (Reactant); RACT (Reactant or reagent) (asym. dihydroxylation of, chiral dihydroquinidine derivative for osmium catalyzed) 100-42-5, reactions 103-30-0, trans-Stilbene 98-83-9, reactions 766-90-5 771-98-2, 1-Phenylcyclohexene 827-54-3, 2-Vinylnaphthalene 872-05-9, 1-Decene 873-66-5 1463-04-3 3901-07-3, trans-Methyl 6714-96-1 7433-56-9, trans-5-Deceme 4-methoxycinnamate 13269-52-8, trans-3-Hexene 13633-26-6 17343-88-3 61142-41-4, Vinylcyclooctane RL: RCT (Reactant); RACT (Reactant or reagent) (asym. dihydroxylation of, chiral dihydroquinidine ligand for osmium catalyzed) 56-54-2, Quinidine RL: RCT (Reactant); RACT (Reactant or reagent) (hydrogenation of) 2325-10-2P 16355-00-3P 25779-13-9P 31612-63-2P 32345-64-5P 34281-90-8P 35638-92-7P 40421-51-0P 40560-98-3P 49801-14-1P 52340-78-0P 53448-10-5P 57495-92-8P 77977-74-3P 78805-31-9P 79299-22-2P 83603-02-5P 84276-14-2P 84518-30-9P 87827-60-9P 87858-06-8P 88196-06-9P 89063-88-7P 98464-24-5P 99881-77-3P 108666-29-1P 108741-12-4P 110549-79-6P 113162-03-1P 113162-04-2P 113162-11-1P 115889-27-5P 121564-12-3P 122517-80-0P 124649-67-8P 125132-75-4P 130876-03-8P 130932-13-7P 130932-14-8P 132486-47-6P 135042-86-3P 135042-87-4P 135096-77-4P 136031-93-1P 138890-41-2P 139093-40-6P 139093-41-7P 139093-42-8P 139093-44-0P 139093-45-1P 139093-46-2P 139093-47-3P 139165-53-0P 139165-54-1P 139165-55-2P 139165-56-3P 139165-57-4P 139165-58-5P 139165-59-6P 139165-61-0P 139165-62-1P 139165-63-2P 139165-64-3P 139165-60-9P 139165-65-4P 139165-66-5P 139165-67-6P 139165-68-7P 139165-69-8P 139165-70-1P 143536-11-2P 149519-66-4P 149519-67-5P 149519-68-6P 149519-69-7P 149562-10-7P 149562-11-8P RL: SPN (Synthetic preparation); PPEP (Preparation) (preparation of) 72989-10-7P 120385-14-0P 139093-52-0P 146333-99-5P 146334-00-1P 146334-01-2P 146334-02-3P 146334-03-4P 146334-04-5P 146334-05-6P

146334-08-9P

146334-09-0P

146334-10-3P

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146334-06-7P

146334-07-8P

149519-70-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as ligand for osmium catalyzed asym. dihydroxylation)

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as ligand for osmium tetroxide catalyzed dihydroxylation of olefins)

=> d 19 ibib abs 6-10

L9 ANSWER 6 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1993:169109 CAPLUS Full-text

DOCUMENT NUMBER: 118:169109

ORIGINAL REFERENCE NO.: 118:29012h,29013a TITLE: Preparation of

(tetrazolylbiphenylmethyl)benzazepinones and related

compounds as growth hormone release promoters $% \left(x\right) =\left(x\right) +\left(x$

INVENTOR(S): Fisher, Michael H.; Wyvratt, Matthew J.; Schoen,

William R.; Devita, Robert J.

PATENT ASSIGNEE(S): Merck and Co., Inc., USA SOURCE: PCT Int. Appl., 346 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	IENT NO.			KIND DATE			APPLICATION NO. DATE	
WO				A1			WO 1992-US2271 19920319 PL, RO, RU, SD	<
US	5206235						US 1992-839742 19920228	<
EP	513974						EP 1992-302143 19920312	
EP	513974			В1		19960904		
	R: AT,	BE,	CH,	DE,	DK.	, ES, FR,	GB, GR, IT, LI, LU, NL, PT, SE	
AT	142206			T		19960915	AT 1992-302143 19920312	<
IL	101206			A		19970218	IL 1992-101206 19920312	<
CA	2063185			A1		19920921	CA 1992-2063185 19920317	<
AU	9213012			Α		19920924	AU 1992-13012 19920319	<
AU	653992			В2		19941020		
CN	1066070			Α		19921111	CN 1992-102954 19920319	<
CN	1033584			С		19961218		
ZA	9202009			A		19921125	ZA 1992-2009 19920319	<
JP	06172316			A		19940621	JP 1992-112069 19920319	<
JP	08000814			В		19960110		
HU	66796			A2		19941228	HU 1992-915 19920319	<
RO	117326			В1		20020130	RO 1993-1245 19920319	<
US	5310737			A		19940510	US 1993-12190 19930202	<
PRIORIT	Y APPLN.	INFO.	. :				US 1991-673695 A 19910320	
							US 1992-839742 A 19920228	<
							WO 1992-US2271 W 19920319	<

OTHER SOURCE(S): MARPAT 118:169109

GI

AΒ Title compds. [I; L = (substituted) phenylene; n, w = 0, 1; p = 0-3; q = 0-4; X = CO, O, S, SO, SO2, CH(OH), CH:CH, imino; R1, R2, R7, R8 = H, halo, (perfluoro)alkyl, perfluoroalkoxy, cyano, NO2, (substituted) Ph, acyl(alkyl), etc.; R4, R5 = H, (substituted) Ph, alkyl, alkenyl, alkynyl, alkanoyloxy, alkoxycarbonyl, carboxy, CHO, amino; R4R5 = (CH2)rB(CH2)s; B = CH2, O, imino, S, SO, SO2; r, s = 1-3; R6 = H, alkyl, Ph, phenylalkyl; R9 = H, (substituted) tetrazolyl, acylalkyl, aminoalkyl, carbamoylalkyl, tetrazolylalkyl, tetrazolylphenyl, tetrazolylphenoxy, etc.; A = (CH2)xCR10R11(CH2)y; x, y = 0-3; R10, R11 = H, CF3, (substituted) alkyl, Ph, etc.; R10R11 = (CH2)t; t = 2-6; R10, R11 may be joined to R4 and/or R5], were prepared for promotion of release of growth hormone (no data). Thus, 3-benzyloxycarbonylamino-3-methyl-N-[2,3,4,5-tetrahydro-2-oxo-1H-1- benzazepin-3R-y1]butanamide (preparation from 3-azido-2,3,4,5-tetrahydro-1H-1-benzazepin-2-one given) was stirred 15 min with NaH in DMF; N-triphenylmethyl-5-[2-(4'-bromobiphen-4-yl)]tetrazole (preparation starting from 5-phenyl-2-trityltetrazole and 4-IC6H4Me given) in DMF was added and the mixture was stirred 90 min to give 95% coupling product, which was bydrogenated in MeOH over Pd(OH)2/C for 14 h to give 89% 3-amino-3methyl-N-[2,3,4,5-tetrahydro-2-oxo-1-[[2'- (1H-tetrazol-5-yl)[1,1'-biphenyl]-4-y1]methy1]-1H-1-benzazepin-3R- y1]butanamide trifluoroacetate.

L9 ANSWER 7 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1990:75971 CAPLUS Full-text

DOCUMENT NUMBER: 112:75971

ORIGINAL REFERENCE NO.: 112:12975a, 12978a

TITLE: Ligand-accelerated catalytic asymmetric

dihydroxylation

INVENTOR(S): Marko, Istvan E.; Sharpless, K. Barry
PATENT ASSIGNEE(S): Massachusetts Institute of Technology, USA

SOURCE: PCT Int. Appl., 66 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

PA:	TENT NO.		KIND		DATE		APPLICATION NO.						
WO	8906225 W: JP			A1		1989	0713	WO	1989-	US86	 -	19890110	<
	RW: AT,	BE,	CH,	DE,	FR,	GB,	ΙΤ,	LU, NI	L, SE				
US	4871855			Α		1989	1003	US	1988-	159068		19880223	<
US	4965364			А		1990	1023	US	1988-	250378		19880928	<
EP	395729			A1		1990	1107	EP	1989-	901900		19890110	<
EP	395729			В1		1995	0927						
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JP	03503885			T		1991	0829	JP	1989-	501814		19890110	<
JP	3153540			В2		2001	0409						
EP	658532			A1		1995	0621	EP	1995-	200458		19890110	<
EP	658532			В1		1999	0407						
	R: AT,	BE,	CH,	DE,	FR,	GB,	ΙΤ,	LI, LU	J, NL,	SE			
AT	128449			T		1995	1015	AT	1989-	901900		19890110	<
AT	178578			T		1999	0415	AT	1995-	200458		19890110	<
JP	20011923	83		Α		2001	0717	JP	2000-	335775		19890110	<
CA	1338314			С		1996	0507	CA	1989-	587964		19890111	<
PRIORIT	Y APPLN.	INFO	.:					US	1988-	142692	А	19880111	<

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US 1988-159068 A 19880223 <--
US 1988-250378
                A 19880928 <--
EP 1989-901900
                 A3 19890110 <--
JP 1989-501814
                 A3 19890110 <--
WO 1989-US86
                 W 19890110 <--
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AΒ Vicinal diols are prepared by asym. dihydroxylation of olefins in the presence of a chiral ligand, an Os-containing catalyst, an amine oxide, an organic solvent, and H2O. OsO4 was injected into a solution of trans-stilbene, hydroquinidine p-chlorobenzoate, and N-methylmorpholine N-oxide in Me2CO and H2O at $0-4^{\circ}$ with shaking, Na2S2O5 added, and the mixture worked up to give 55% (R,R)-PhCH(OH)CH(OH)Ph of 99% enantiomeric excess. Preparation of dihydroquinidine derivs. and their recovery were also given. Asym. dihydroxylations of trans-3-hexene, 1-phenylcyclohexene, β -methylstyrene, and Me (E)-4-methoxycinnamate were also given.

ANSWER 8 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN 1986:129936 CAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 104:129936

ORIGINAL REFERENCE NO.: 104:20565a,20568a

1,5-Benzothiazepine derivatives

INVENTOR(S): Takeda, Mikio; Ohishi, Tokuro; Nakajima, Hiromichi;

Nagao, Taku

Tanabe Seiyaku Co., Ltd. , Japan PATENT ASSIGNEE(S):

Eur. Pat. Appl., 69 pp. SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	PATENT NO.			KIND		DATE		API	PLICATION NO.	DATE	
	158339 158339			A2 A3		19851016 19860611		EP	1985-104341	 19850410	<
	158339			B1		19890118					
	R: AT,	BE,	CH,	DE,	FR,	, GB, IT,	LI,	LU	J, NL, SE		
US	4585768			A		19860429		US	1985-715116	19850322	<
AU	8540392			A		19851017		AU	1985-40392	19850326	<
AU	572978			B2		19880519					
JP	60226866			A		19851112		JΡ	1985-73356	19850405	<
JP	03074659			В		19911127					
DK	8501577			A		19851011		DK	1985-1577	19850409	<
DK	172257			В1		19980209					
HU	37411			A2		19851228		HU	1985-1303	19850409	<
HU	191785			В		19870428					
CA	1218991			A1		19870310		CA	1985-478613	19850409	<
SU	1632372			А3		19910228		SU	1985-3880298	19850409	<
AT	40126			T		19890215		ΑT	1985-104341	19850410	<
CN	85103524			А		19861105		CN	1985-103524	19850502	<
	1017706			В		19920805					
	1358784			A3		19871207		SU	1986-4027451	19860513	<
	Y APPLN.		. :			,			1984-9258		
			•						1985-104341	19850410	
	OHD OF (0)			0-0-		·			1303 101311	13000110	•

OTHER SOURCE(S): CASREACT 104:129936; MARPAT 104:129936

GΙ

AB (Aminoethyl)benzothiazepinones I (R1, R3 = alkyl; R2 = H, alkanoyl, PhCH2; 1 of R4, R5 = C1, the other H; R6 = H) were prepared Thus, 5,2-C1(O2N)C6H3SH and Me trans-3-(4-methoxyphenyl)oxiranecarboxylate were stirred in PhMe containing Zn(OAc)2 to give (±)-threo-5,2-C1C6H3SCH(C6H4OMe-4)CH(OH)CO2Me. This was saponified and the acid resolved by crystallization of its L-lysine salt. The (-)-threo isomer was hydrogenated over Pd/C to give the amine which was cyclized by refluxing in xylene to give (-)-cis-I (R2 = R3 = Me, R2 = R5 = H, R4 = C1, R6 = PhCH2) which was treated successively with PhCH2O2CCl and HBr/HOAc to give (-)-cis-I (R1 = R3 = Me, R2 = R5 = R6 = N, R4 = C1), isolated as its hydrochloride (II). Blood plasma from rats given 10 mg II/kg orally showed ≥50% inhibition of collagen-induced platelet aggregation.

L9 ANSWER 9 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1976:58822 CAPLUS Full-text

DOCUMENT NUMBER: 84:58822

ORIGINAL REFERENCE NO.: 84:9659a,9662a

TITLE: Synthesis of quaiacylglycerol- β -quaiacyl ether

AUTHOR(S): Nakatsubo, Fumiaki; Sato, Kimihiko; Higuchi, Takayoshi

CORPORATE SOURCE: Wood Res. Inst., Kyoto Univ., Uji, Japan

SOURCE: Holzforschung (1975), 29(5), 165-8

CODEN: HOLZAZ; ISSN: 0018-3830

DOCUMENT TYPE: Journal LANGUAGE: English

GI For diagram(s), see printed CA Issue.

Guaiacylglycerol- β -guaiacyl ether (I), the model compound of arylglycerol- β -aryl ether structure in lignin was synthesized in high yield through five reaction steps from vanillin. The key step of this synthetic method was the condensation reaction between o-MeOC6H4OCH2CO2Et and 3,4-(MeO) (BzO)C6H3CHO (II). At this step, lithium diisopropyl amide was used as the base, and 3,4-(MeO) (BzO)C6H3CH(OH)CH(CO2Et)OC6H4OMe-o (III) was obtained in 95% yield as an oily substance consisted of two isomers, from which only erythro isomer was obtained as crystal in 51% yield. The residual oily substance was converted to its carbamate (IV) and crystallized in 70% yield. The crystalline III and IV were then converted to I by the LiAlH4 reduction and subsequent hydrogenation with Pd-C. The overall yield of I from II was about 72%.

L9 ANSWER 10 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1975:443121 CAPLUS Full-text

DOCUMENT NUMBER: 83:43121

ORIGINAL REFERENCE NO.: 83:6811a,6814a

TITLE: Polyphenolic acids of Lithospermum ruderale (Boraginaceae). I. Isolation and structure

determination of lithospermic acid

AUTHOR(S): Kelley, Charles J.; Mahajan, J. R.; Brooks, Lucille C.; Neubert, Leonard A.; Breneman, W. R.; Carmack,

Marvin

CORPORATE SOURCE: Dep. Chem., Indiana Univ., Bloomington, IN, USA SOURCE: Journal of Organic Chemistry (1978), 40(12),

1804-15

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

GI For diagram(s), see printed CA Issue.

AB A structure is proposed for lithospermic acid (I), C27H22O12, the major polyphenolic acid of Lithospermum ruderale and several other plant species of the families, Boraginaceae and Labiatae. Chromatog. on Sephadex of aqueous

exts. of the plant yields the di-K salt of I, together with salts of lesser constituents which include (R)-3-(3,4-dihydroxyphenyl)lactic acid, 2-(3,4-dihydroxyphenyl)-3-carboxy-4-(2-carboxy-trans-vinyl)-7- hydroxycoumaran, and rosmarinic acid. Structures were deduced from spectral studies of the salts, the free acids, and also the methylated derivs., produced by the action of CH2N2 on the free acids or Me2SO4 on the salts.

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L9 ANSWER 11 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1966:51892 CAPLUS Full-text

DOCUMENT NUMBER: 64:51892

ORIGINAL REFERENCE NO.: 64:9672d-h,9673a-c

TITLE: New synthesis of 2-hydroxy-2-benzyl-3-coumaranones AUTHOR(S): Chopin, Jean; Durual, Pierre; Chadenson, Michele

CORPORATE SOURCE: Fac. Sci., Lyon

SOURCE: Bulletin de la Societe Chimique de France (

1965), (12), 3572-7

CODEN: BSCFAS; ISSN: 0037-8968

DOCUMENT TYPE: Journal LANGUAGE: French

OTHER SOURCE(S): CASREACT 64:51892

A series of 2-hydroxy-2-benzyl-3-coumaranones was prepared by the AΒ debenzylation of the appropriate α -diketones obtained by the isomerization of 2'-benzyloxychalcones or of the corresponding 2'-benzyloxy- α -methoxychalcones. The existence of an equilibrium between the cyclic and open forms was demonstrated by their N. M.R. spectra and was in good agreement with the results of the alkaline rearrangement of the 3-hydroxyflavanones. 2,4-(HO) 2C6H3COCH2OMe (I) (5.5 q.) with Me2SO4 yielded 4.1 q. 2,4-HO(MeO)C6H3COCH2OMe (II), m. 65-6° (EtOH). II (1.9 g.) with PhCH2Cl in HCONMe2 in the presence of NaI and K2CO3 gave 1.8 g. 2,4phCH2O(MeO)C6H3COCH2OMe (III), m. 66° (EtOH). I (2 g.) with PhCH2Cl gave similarly 2.6 g. 2,4-(PhCH2O)2C6H3COCH2OMe (IV), m. 104° (EtOH). II (1 g.) and 1 g. BzH in 20 cc. EtOH treated overnight with 2 g. 50% ag. NaOH and acidified yielded 785 mg. 2,4-PhCH2O(MeO)C6H3COC(OMe):CHPh (V), m. 95° (EtOH). V (1 q.) in 100 cc. MeOH and 20 cc. H2O refluxed 6 hrs. with 10 cc. concentrated HCl gave 550 mg. yellow 2,4-PhCH2O(MeO)C6H3COCOCH2Ph, m. 86° (EtOH). IV (1 g.) with 1 g. BzH gave 1.1 g. yellowish white 2,4-(PhCH2O)2-C6H3COC(OMe): CHPh (VI); m. 120°. III (1 g.) and 1 g. p-MeOC6H4CHO yielded 815 mg. yellowish white 4-MeO analog of VI, m. 92-3°. o-PhCH2OC6H4COCOCH2Ph (1 g.) in 20 cc. AcOH heated 1.5 hrs. on a water bath with 10 cc. concentrated HCl yielded 585 mg. beige 2-hydroxy-2-benzyl-3-coumaranone (VII), m. 104° (C6H6-hexane). o-PhCH2OC6H4COCH(OH)CHClPh gave similarly 78% VII. o-PhCH2OC6H4COCOCH2C6H4OMe-p (1 g.) gave similarly 490 mg. beige 4'-MeO derivative (VIII) of VII, m. 120° (C6H6-hexane). 3-Hydroxy-4'-methoxyflavanone (IX) (1 q.) heated 5 min. on the water bath with 100 cc. 2N alc. KOH and poured into 200 cc. 2N HCl gave 403 mg. 4'-methoxyflavonol (X), m. 228° (EtOH); the filtrate from the X yielded 385 mg. oily o-HOC6H4C(OH)(CO2H)CH2C6H4OMe-p (XI) which gave an intense blue color with alc. FeCl3. The XI methylated with Me2SO4 and K2CO3 in MeOH gave o-MeOC6H4C(OH)(CO2Me)CH2C6H4OMe-p, m. 133° (EtOH), which saponified with alc. KOH yielded o-MeOC6H4C(OH)(CO2H)CH6C6H4OMe-p, m. 159° (EtOH), and 190 mg. VIII, m. 120°. VIII dehydrated with concentrated H2SO4 gave 4'-methoxyaurone, m. 138-9°. IX heated 15 min. on the water bath with alc. KOH gave 646 mg. XI and 296 mg. X. 4,2-MeO(PhCH2O)C6H3COCOCH2Ph (XII) (1 g.) with HCl-AcOH gave 435 mg. 6-MeO derivative (XIII) of VII, m. 120° (C6H6-hexane). XII (1 g.) in 20 cc. EtOH hydrogenated over 100 mg. 10% Pd-C, and the product

chromatographed on Al2O3 yielded 450 mg. XIII. 4,2-MeO(PhCH2O)C6H3COCH(OH)CH-ClPh with HCl-AcOH gave 55% XIII. V gave similarly 57% XIII. XIII (100 mg.) and 2 cc. concentrated H2SO4 heated 10 min. on the water bath gave 76 mg. 6methoxyaurone, m. 145° (EtOH). XIII (100 mg.) in 5 cc. EtOH and 3 cc. 2N KOH heated 3 min. on the water bath and acidified with 2N HCl yielded oily 2,4-HO(MeO)C6H3C(OH)(CO2H)CH2Ph (it gave an intense blue color with alc. FeCl3) which heated 3 min. on the water bath gave 79 mg. 6-methoxy-3-benzal-2coumaranone, m. 129° (MeOH). 4,-2-MeO(PhCH2O)C6H3COCOCH2C6H4OMe-p (1 g.) with HClAcOH vielded 470 mg. beige 4',6-dimethoxy derivative (XIV) of VII, m. 111° (C6H6-hexane). 4,2-MeO(PhCH2O)C6H3COC(OMe):CHC6H4OMe-p (500 mg.) gave similarly 225 mg. XIV. XIV treated with concentrated H2SO4 gave 6,4'dimethoxyaurone, m. 134°. 2,4-(PhCH2O)2C6H3COC(OMe):CHPh (500 mg.) with HClAcOH gave 179 mg. 6-PhCH2 derivative of VII, m. 186-7° (C6H6), which dehydrated with concentrated H2SO4 yielded 6-hydroxyaurone, m. 262-5°. 2'-Hydroxy-3,4-dimethoxychalcone (8 g.) with Ac2O and AcONa gave 8 g. acetate, m. 90°, which treated in 100 cc. CS2 and 10 cc. CH2Cl2 with 1.25 cc. Br in 10 cc. CS2 and kept 1 hr. yielded 11 g. dibromide (XV), m. 162-3° (CHCl3-hexane). XV (11 g.) refluxed 15 min. with 80 cc. Me2CO and 20 cc. H2O and heated 5 min. with 10 g. Na2CO3 in 70 cc. H2O gave 1.8 g. 3',4'-dimethoxyflavanol (XVI), m. $156-8^{\circ}$ (MeOH). 3-Hydroxy-3',4'-dimethoxyflavanone (XVII) heated 5 min. on the water bath with 2N alc. KOH gave 235 mg. 3-hydroxy-3',4'-dimethoxyflavone, m. $196-7^{\circ}$, 275 mg. o-HOC6H4C(OH)(CO2H)CH2C6H4(OMe)2-3,4 (XVIII) (it gave an intense blue color with alc. FeCl3), and 365 mg. 3',4'-dimethoxy derivative of VII. XVIII with Me2SO4 yielded o-MeOC6H4C(OH)(CO2Me)CH2C6H3(OMe)2-3,4, m. 129° (EtOH), which saponified with alc. KOH gave o-MeOC6H4C(OH)(CO2H)CH2C6H3(OMe)2-3,4, m. 179° (EtOH). XVII gave similarly during 15 min. 45% XVIII and 45% XVI.

TI New synthesis of 2-hydroxy-2-benzyl-3-coumaranones

SO Bulletin de la Societe Chimique de France (1965), (12), 3572-7

L9 ANSWER 12 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1965:438715 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 63:38715
ORIGINAL REFERENCE NO.: 63:6873c-g

TITLE: Synthesis of prephenic acid diethyl acetal and its

hydrolysis to phenylpyruvic acid and prephenic acid

AUTHOR(S): Plieninger, Hans; Arnold, Lothar; Fischer, Rolf;

Hoffmann, Werner

CORPORATE SOURCE: Univ. Heidelberg, Germany

SOURCE: Chemische Berichte (1965), 98(6), 1774-81

CODEN: CHBEAM; ISSN: 0009-2940

DOCUMENT TYPE: Journal LANGUAGE: German

GI For diagram(s), see printed CA Issue.

AB I and the di-Et acetal (II) of III were prepared The time at which the maximum yield of III can be obtained by the acid hydrolysis of II was calculated from kinetic data and the calcn. confirmed by the experiment; III was formed together with larger amts. of PhCH2COCO2H (IV). Di-Et 2cyclohexen-4-one-1-carboxylate-1-pyruvate di-Et acetal (35.6 g.), b0.1 155°, n25D 1.4087, in 350 cc. tert-BuOH refluxed 5 hrs. with stirring with 5 cc. AcOH and 11 g. SeO2, treated with an addnl. 11 g. SeO2, and again refluxed 5 hrs., and the product shaken in Et2O 6 hrs. with 10 g. deactivated Raney Ni yielded 25 g. I, b0.1 150°, n20D 1.4840. I (35.6 mg.), 5.0 cc. EtOH, and 4.0cc. 0.1N NaOH heated 0.5 hr. at 50° , treated with 5 cc. N HCl, and heated 15 min. at 50° , and the mixture cooled and diluted with H2O to 100 cc. gave a solution containing 18 mg. p-HOC6H4CH2COCO2H. I (1.0 g.), 100 mg. Na, and 5 cc. EtOH heated 3 hrs. at 50° gave 0.70 g. p-HOC6H4CH24C(OEt)2CO2Et (IV). I (7.1 g.) in 10 cc. EtOH added dropwise with stirring during 15 min. at 5° to 0.40 g. NaBH4 in 40 cc. EtOH and kept 15 min. at 20° yielded 5.3 g. oily di-Et ester (V) of II, n25D 1.4665, and 1.1 g. IV. V (3.6 g.) in 30 cc. EtOH treated 2 days at room temperature with 1.6 g. NaOH in 20 cc. H2O and evaporated to 15 cc., buffered to pH 8 with N HCl, and diluted with H2O to 25 cc. gave a 0.4M solution; a 0.1-cc. portion and 4 cc. N HCl heated 15 min. at 50° and the mixture adjusted to pH 14 with N NaOH gave a 0.3M II solution; a 6.25-cc. portion stirred with 4.25 g. Ba(OAc)2, centrifuged to remove some solid, diluted with about 200 cc. EtOH, and kept several hrs. yielded 1.3 g. Ba salt; a 0.5-g. portion in 10 cc. H2O hydrogenated over PdBaSO4, filtered, and treated 20 hrs. with 2,4-(O2N)2C6H3NHNH2 in 2N HCl, and the precipitate chromatographed on paper showed the presence of the 2,4-dinitrophenylhydrazones of cis- and trans-tetrahydroprephenic acid (relative to OH and CO2H groups). II (0.3M aqueous solution) (10 cc.) adjusted with N HCl to pH 1.8, kept 10 min. at 20°, neutralized with N NaOH, and treated with 1.0 g. Ba(OAc)2 in 5 cc, H2O gave 0.5 g. Ba salt of a 6:50 mole ratio mixture of III and IV.

- TI Synthesis of prephenic acid diethyl acetal and its hydrolysis to phenylpyruvic acid and prephenic acid
- SO Chemische Berichte (1965), 98(6), 1774-81 CODEN: CHBEAM; ISSN: 0009-2940
- I and the di-Et acetal (II) of III were prepared The time at which the AΒ maximum yield of III can be obtained by the acid hydrolysis of II was calculated from kinetic data and the calcn. confirmed by the experiment; III was formed together with larger amts. of PhCH2COCO2H (IV). Di-Et 2cyclohexen-4-one-1-carboxylate-1-pyruvate di-Et acetal (35.6 g.), b0.1 155°, n25D 1.4087, in 350 cc. tert-BuOH refluxed 5 hrs. with stirring with 5 cc. AcOH and 11 g. SeO2, treated with an addnl. 11 g. SeO2, and again refluxed 5 hrs., and the product shaken in Et2O 6 hrs. with 10 g. deactivated Raney Ni yielded 25 g. I, b0.1 150°, n20D 1.4840. I (35.6 mg.), 5.0 cc. EtOH, and 4.0cc. 0.1N NaOH heated 0.5 hr. at 50°, treated with 5 cc. N HCl, and heated 15 min. at 50° , and the mixture cooled and diluted with H2O to 100 cc. gave a solution containing 18 mg. p-HOC6H4CH2COCO2H. I (1.0 g.), 100 mg. Na, and 5 cc. EtOH heated 3 hrs. at 50° gave 0.70 g. p-HOC6H4CH24C(OEt)2CO2Et (IV). (7.1 g.) in 10 cc. EtOH added dropwise with stirring during 15 min. at 5° to 0.40 g. NaBH4 in 40 cc. EtOH and kept 15 min. at 20° yielded 5.3 g. oily di-Et ester (V) of II, n25D 1.4665, and 1.1 g. IV. V (3.6 g.) in 30 cc. EtOH treated 2 days at room temperature with 1.6 g. NaOH in 20 cc. H2O and evaporated to 15 cc., buffered to pH 8 with N HCl, and diluted with H2O to 25 cc. gave a $0.4 \mathrm{M}$ solution; a $0.1 - \mathrm{cc}$. portion and $4 \mathrm{cc}$. N HCl heated 15 min. at 50° and the mixture adjusted to pH 14 with N NaOH gave a 0.3M II solution; a 6.25-cc. portion stirred with 4.25 g. Ba(OAc)2, centrifuged to remove some solid, diluted with about 200 cc. EtOH, and kept several hrs. yielded 1.3 g. Ba salt; a 0.5-q. portion in 10 cc. H2O hydrogenated over PdBaSO4, filtered, and treated 20 hrs. with 2,4-(O2N)2C6H3NHNH2 in 2N HCl, and the precipitate chromatographed on paper showed the presence of the 2,4dinitrophenylhydrazones of cis- and trans-tetrahydroprephenic acid (relative to OH and CO2H groups). II (0.3M aqueous solution) (10 cc.) adjusted with N HCl to pH 1.8, kept 10 min. at 20°, neutralized with N NaOH, and treated with 1.0 g. Ba(OAc)2 in 5 cc, H2O gave 0.5 g. Ba salt of a 6:50 mole ratio mixture of III and IV.
- 126-49-8P, 2,5-Cyclohexadiene-1-pyruvic acid, 1-carboxy-4-hydroxy-156-06-9P, Pyruvic acid, phenyl- 156-39-8P, Pyruvic acid, (p-hydroxyphenyl)- 2931-07-9P, 2,5-Cyclohexadiene-1-propionic acid, 1-carboxy-α,α-diethoxy-4-hydroxy-, diethyl ester 2931-07-9P, 2,5-Cyclohexadiene-1-pyruvic acid, 1-carboxy-4-hydroxy-, diethyl ester, di-Et acetal 2931-35-3P, 2,5-Cyclohexadiene-1-pyruvic acid, 1-carboxy-4-hydroxy-, diethyl acetal 2931-35-3P, 2,5-Cyclohexadiene-1-propionic acid, 1-carboxy-α,α-diethoxy-4-hydroxy- 2931-38-6P, 2-Cyclohexene-1-propionic acid, 1-carboxy-α,α-diethoxy-4-oxo-,

diethyl ester 2931-38-6P, 2-Cyclohexene-1-pyruvic acid, 1-carboxy-4-oxo-, diethyl ester, di-Et acetal 2931-39-7P, 2,5-Cyclohexadiene-1-propionic acid, 1-carboxy-α,α-diethoxy-4-oxo-, diethyl ester 4375-92-2P, Pyruvic acid, (p-hydroxyphenyl)-, ethyl ester, di-Et acetal 4387-06-8P, 1,4-Methanonaphthalene-5,8,9-trione, 1-bromo-1,4,4a,8a-tetrahydro-, 9-(dimethyl acetal) 4387-08-0P, 1,4-Methanonaphthalene-5,8-dione, 9,9-diethoxy-1,4,4a,8a-tetrahydro-4423-91-0P, 5-Norbornene-2,3-dicarboxylic anhydride, 7-oxo-825632-90-4P, 2,5-Cyclohexadiene-1-pyruvic acid, 1-carboxy-4-oxo-, diethyl ester, di-Et acetal RL: PPEP (Preparation) (preparation of)

L9 ANSWER 13 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1963:435348 CAPLUS Full-text

DOCUMENT NUMBER: 59:35348

ORIGINAL REFERENCE NO.: 59:6301g-h,6302a-d

TITLE: Preparation of the threo- and erythro-forms of

DL-guaiacylglycerol and of DL-veratrylglycerol

AUTHOR(S): Adler, Erich; Gustafsson, Bo

CORPORATE SOURCE: Chalmers Tek. Hogskola, Goteborg, Swed. SOURCE: Acta Chemica Scandinavica (1963), 17, 27-36

CODEN: ACHSE7; ISSN: 0904-213X

DOCUMENT TYPE: Journal LANGUAGE: German

OTHER SOURCE(S): CASREACT 59:35348

aB cf. CA 48, 5147g. Hydrogenation of 3,4-dimethoxyphenylpropiolic acid in EtOH with Lindlar catalyst 50 min. gave 70% cis-methylferulic acid (I), m. 104°, which with CH2N2 yielded 90% Me ester, prisms, m. 92-3°. Reduction of 30 g. Me trans-methylferulate (II) in 200 cc. dioxane with 5.7 g. LiAlH4 in 500 cc. absolute Et2O, addition of H2O and dilute H2SO4, extraction with CHC18, and distillation of the residue of the CHC13 extract gave 70% methylconiferyl alc., b5 110-20°, needles, m. 79-80°; Ac derivative (III) b12 190-5°. Treating 80 mg. III in 3 cc. Et2O-C5H5N (25:1) 16 hrs. with 0.1 g. 0sO4 in 2 cc. Et2O and hydrolyzing the osmic ester in 2 cc. EtOH with 0.65 g. Na2SO3 in 3 cc. H2O 1 hr. at 100°, evaporating the filtered solution in vacuo, and extracting the residue with CHC13 gave 85% DL-threoveratrylglycerol (IV), m. 109-10°. Treating II similarly with OsO4 and acetylating the hydrolyzed Os ester gave 60%

DL-threo- α , β -diacetoxymethylhydroferulic acid (V), prisms, m. 148-9°. Esterification of V with CH2N2 followed by reduction yielded IV. Refluxing 15 g. Me α -bromo- β -acetoxymethyl-hydroferulate in 65 cc. AcOH and 65 cc. Ac2O 45 min. with 7 q. AgOAc, treating the filtered solution with H2O, evaporating the solution in vacuo, extracting the residue with CHC13, evaporating the washed (NaHCO3) solution, and crystallizing the residue from Et20-C6H14 gave 41% Me threo- α , β -diacetoxymethylhydroferulate (VI), m. 102-3°. Concentration of the mother liquor gave 40% erythro isomer (VII), prisms, m. 71-3°. VI was also obtained in 10% yield when 4.44 g. II was oxidized with KMnO4 at $-50\,^{\circ}$ according to Riiber (CA 9, 2244). Reduction of VII with LiAlH4 gave DLerythroveratrylglycerol (VIII), plates, m. 92-3°, λ maximum 278 m μ (log ϵ 3.52). The infrared spectra of IV and VIII differed distinctly. When Me erythro-diacetoxyacetylhydroferulate was reduced with LiAlH4 (cf. loc. cit.) and the acid step was avoided by neutralizing the reaction mixture with AcOH, 55% DL-erythro-guaiacylglycerol (IX), m. $83-4^{\circ}$ was obtained; IX tetraacetate m. $86-8^{\circ}$. IX and CH2N2 gave 90% VIII. Treating 0.31 g. IX with 0.1N H2SO4 neutralizing the mixture with BaCO8, evaporating the filtered solution in vacuo, and treating the residue with moist EtOAc gave 0.18 g. unchanged IX and, from the mother liquor, 0.05 g. threo-DL-guaiacylglycerol (X);

tetraacetate m. 113-14°. Benzyl-coniferyl alc. benzoate (0.515 g.) (Freudenberg and Achtzehn, CA 50, 1661h) was treated in 12 cc. Et20-C5H5N with 0.35 g. 0s04 and the precipitate formed was boiled 1 hr. with 2.3 g. Na2SO3 in 10 cc. H2O, giving threo-(O-benzylguaiacyl)glycerol, m. 101°, which when treated in EtOH with prehydrogenated PdC12-BaSO4, yielded X as a sirup; tetraacetate, 70%, m. 113-14°. X and CH2N2 gave 70% IV. Careful fractionation of the mother liquor of the Me erythro- α , β -diacetoxyacetylhydroferulate gave 10% threo isomer, prisms, m. 81-2°, which with LiAlH4 gave 55% X. Treating 0.074 g. trans-methylisoeugenol (XI) with 0s04 gave 75% DL-threo-methylisoeugenol glycol, m. 90°, which was also obtained in 15% yield when 1.93 g. XI was treated at -50° with 1.96 g. KMnO4. Acetylation of 3.3 g. XI in 3 cc. AcOH with 6.6 g. Pb(OAc)4 and reduction of the α , β -diacetoxyveratrylpropane formed with LiAlH4 gave 2 g. of a product, m. 80-100°, which on fractional crystallization yielded crythro-methylisoeugenol glycol, m. 123°, and the threo isomer, m. 88°.

- $\mbox{{\tt TI}}\mbox{{\tt Preparation}}$ of the threo- and erythro-forms of DL-guaiacylglycerol and of DL-veratrylglycerol
- SO Acta Chemica Scandinavica (1963), 17, 27-36 CODEN: ACHSE7; ISSN: 0904-213X
- ab cf. CA 48, 5147g. Hydrogenation of 3,4-dimethoxyphenylpropiolic acid in EtOH with Lindlar catalyst 50 min. gave 70% cis-methylferulic acid (I), m. 104°, which with CH2N2 yielded 90% Me ester, prisms, m. 92-3°. Reduction of 30 g. Me trans-methylferulate (II) in 200 cc. dioxane with 5.7 g. LiAlH4 in 500 cc. absolute Et2O, addition of H2O and dilute H2SO4, extraction with CHC18, and distillation of the residue of the CHC13 extract gave 70% methylconiferyl alc., b5 110-20°, needles, m. 79-80°; Ac derivative (III) b12 190-5°. Treating 80 mg. III in 3 cc. Et2O-C5H5N (25:1) 16 hrs. with 0.1 g. 0sO4 in 2 cc. Et2O and hydrolyzing the osmic ester in 2 cc. EtOH with 0.65 g. Na2SO3 in 3 cc. H2O 1 hr. at 100°, evaporating the filtered solution in vacuo, and extracting the residue with CHC13 gave 85% DL-threoveratrylglycerol (IV), m. 109-10°. Treating II similarly with 0sO4 and acetylating the hydrolyzed Os ester gave 60%

DL-threo- α , β -diacetoxymethylhydroferulic acid (V), prisms, m. 148-9°. Esterification of V with CH2N2 followed by reduction yielded IV. Refluxing 15 g. Me α -bromo- β -acetoxymethyl-hydroferulate in 65 cc. AcOH and 65 cc. Ac2O 45 min. with 7 g. AgOAc, treating the filtered solution with H2O, evaporating the solution in vacuo, extracting the residue with CHC13, evaporating the washed (NaHCO3) solution, and crystallizing the residue from Et20-C6H14 gave 41% Me threo- α , β -diacetoxymethylhydroferulate (VI), m. 102-3°. Concentration of the mother liquor gave 40% erythro isomer (VII), prisms, m. 71-3°. VI was also obtained in 10% yield when 4.44 g. II was oxidized with KMnO4 at -50° according to Riiber (CA 9, 2244). Reduction of VII with LiAlH4 gave DLerythroveratrylglycerol (VIII), plates, m. 92-3°, λ maximum 278 m μ (log ϵ 3.52). The infrared spectra of IV and VIII differed distinctly. When Me erythro-diacetoxyacetylhydroferulate was reduced with LiAlH4 (cf. loc. cit.) and the acid step was avoided by neutralizing the reaction mixture with AcOH, 55% DL-erythro-guaiacylglycerol (IX), m. $83-4^{\circ}$ was obtained; IX tetraacetate m. $86-8^{\circ}$. IX and CH2N2 gave 90% VIII. Treating 0.31 g. IX with 0.1N H2SO4 neutralizing the mixture with BaCO8, evaporating the filtered solution in vacuo, and treating the residue with moist EtOAc gave 0.18 g. unchanged IX and, from the mother liquor, 0.05 q. threo-DL-quaiacylqlycerol (X); tetraacetate m. 113-14°. Benzyl-coniferyl alc. benzoate (0.515 g.) (Freudenberg and Achtzehn, CA 50, 1661h) was treated in 12 cc. Et20-C5H5N with 0.35 g. 0s04 and the precipitate formed was boiled 1 hr. with 2.3 g. Na2S03 in 10 cc. H2O, giving threo-(O-benzylguaiacyl)glycerol, m. 101°, which when treated in EtOH with prehydrogenated PdCl2-BaSO4, yielded X as a sirup; tetraacetate, 70%, m. 113-14°. X and CH2N2 gave 70% IV. Careful fractionation of the mother liquor of the Me erythro- α , β -diacetoxyacetylhydroferulate gave 10% threo isomer, prisms, m. 81-2°, which with LiAlH4 gave 55% X. Treating

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methylisoeugenol glycol, m. 90°, which was also obtained in 15% yield when
     1.93 g. XI was treated at -50^{\circ} with 1.96 g. KMnO4. Acetylation of 3.3 g. XI
     in 3 cc. AcOH with 6.6 q. Pb(OAc)4 and reduction of the \alpha, \beta-
     diacetoxyveratrylpropane formed with LiAlH4 gave 2 g. of a product, m. 80-
     100°, which on fractional crystallization yielded crythro-methylisoeugenol
     glycol, m. 123°, and the three isomer, m. 88°.
     4756-11-0P, 1,2,3-Propanetriol, 1-(3,4-dimethoxyphenyl)-, DL-erythro-
ΙT
     14737-88-3P, Cinnamic acid, 3,4-dimethoxy-, cis- 18523-76-7P,
     2-Propen-1-ol, 3-(3,4-dimethoxyphenyl)-
                                                20133-19-1P, 1,2-Propanediol,
     1-(3,4-dimethoxyphenyl)-, DL-erythro-
                                              20133-19-1P, 1,2-Propanediol,
     1-(3,4-dimethoxyphenyl)-, DL-threo- 27391-16-8P, 1,2,3-Propanetriol,
     1-(4-hydroxy-3-methoxyphenyl)-, DL-threo- 30461-78-0P, Cinnamic acid,
     3,4-dimethoxy-, methyl ester, cis- 38916-91-5P, 1,2,3-Propanetriol,
     1-(4-hydroxy-3-methoxyphenyl)-, DL-erythro- 65401-84-5P, 2-Propen-1-ol,
     3-(3,4-dimethoxyphenyl)-, acetate 69731-37-9P, 1,2,3-Propanetriol,
     1-(4-hydroxy-3-methoxypheny1)-, tetraacetate, DL-erythro- 69731-38-0P,
     1,2,3-Propanetriol, 1-(4-hydroxy-3-methoxyphenyl)-, tetraacetate,
                69731-40-4P, 1,2,3-Propanetriol, 1-(3,4-dimethoxyphenyl)-,
     DL-threo-
     DL-threo-
                 92582-54-2P, Hydrocinnamic acid,
     \alpha, \beta-dihydroxy-3, 4-dimethoxy-, methyl ester, diacetate,
     DL-erythro- 92582-54-2P, Hydrocinnamic acid,
     \alpha, \beta-dihydroxy-3, 4-dimethoxy-, diacetate, DL-threo-
     93160-12-4P, Hydrocinnamic acid,
     \alpha, \beta-dihydroxy-3, 4-dimethoxy-, methyl ester, diacetate,
     DL-threo-
                95560-55-7P, 2-Propen-1-ol,
     3-[4-(benzyloxy)-3-methoxyphenyl]-, benzoate
                                                     876369-20-9P,
     1,2,3-Propanetriol, 1-[4-(benzyloxy)-3-methoxyphenyl]-, threo-
     882025-72-1P, Hydrocinnamic acid,
     \alpha, \beta, 4-trihydroxy-3-methoxy-, methyl ester, triacetate, erythro-
     882025-73-2P, Hydrocinnamic acid,
     \alpha, \beta, 4-trihydroxy-3-methoxy-, methyl ester, threo-
     RL: PREP (Preparation)
        (preparation of)
     ANSWER 14 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                         1958:35300 CAPLUS Full-text
DOCUMENT NUMBER:
                         52:35300
ORIGINAL REFERENCE NO.: 52:6362c-i,6363a-h
TITLE:
                         Clarifying the constitution of xanthocillin-a new
                         antibiotic
                         Hagedorn, I.; Tonjes, H.
AUTHOR(S):
CORPORATE SOURCE:
                         Tech. Hochschule, Dresden, Germany
                         Pharmazie (1957), 12, 567-80
SOURCE:
                         CODEN: PHARAT; ISSN: 0031-7144
DOCUMENT TYPE:
                         Journal
                         Unavailable
LANGUAGE:
     cf. C.A. 50, 17161e. Xanthocillin-X (I), C18H12O2N2, occurs in yellow
AΒ
     crystals, decompose 220°. Di-Me ether of I, m. 181° (decomposition), is
     prepared by treating with CH2N2 in Et2O. Di-Ac derivative, decompose above
     200°, is formed by heating I with Ac2O and anhydrous AcONa to 80° and
     precipitating with H2O. Di-Bz derivative of I, decompose 200°, is formed by
     acetylation with BzCl in NaOH without heating. 1,4-Bis(p-hydroxyphenyl)-2,3-
     butanedione (II), m. 181-2^{\circ}, may be prepared either by heating I with 1:1
     AcOH-HCl at 100^{\circ} 15 min. (yield 8%) or by storing I in 5:3 EtOH-H2SO4 15 hrs.
     at room temperature, treating with H2O, and extracting with Et2O (yield 20%);
     the osazone of II, m. 228-9°. 2,3-Bis(p-hydroxybenzyl)quinoxaline, m. 234-5°
     (decomposition), is prepared by boiling II and o-C6H4(NH2)2 in alc. for a
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0.074 g. trans-methylisoeugenol (XI) with 0s04 gave 75% DL-threo-

short time and precipitating with H2O. 2,3-Bis(p-methoxybenzyl)quinoxaline is prepared from the last compound by treatment with Me2SO4 in NaOH solution (mono-Me ether forms 1st., m. $144-6^{\circ}$; more heat and stronger solution of NaOH gives the di-Me ether, m. 89-90°). 1,4-Bis(p-methoxypheny1)-2,3-butanedione (III), m. $135-6^{\circ}$, is prepared analogously, as in the case of II, but using I di-Me ether in place of I; osazone of III, m. 185-6°; dioxime, m. 201-3°. 2,3-Bis(p-methoxybenzyl)quinoxaline, m. 90-1, is prepared by treating III with o-C6H4(NH2)2. Oxidation of III with CrO3 gave p-MeOC6H4CO2H, m. 182-3°. Oxidation with H2O2 furnished p-MeOC6H4CH2CO2H, m. 85°. When III was heated in AcOH with Zn dust and H2O, the α -hydroxyketone (IV), m. 77-7.5°, was formed. 1,4-Bis(p-methoxyphenyl)-2,3-butanediol (V), m. 160.5-161.50, was formed when LiBH4 in absolute alc. and III in absolute Et20 were mixed and heated 60 min. at 100° (protected from H2O) and the resulting complex destroyed with 2N NaOH (redistn. with NaBH4 led to same product). (AcO)4Pb was added to V at $40-5^{\circ}$, after 30 min. the mixture filtered, the filtrate concentrated, and semicarbazide, HCl, and AcONa added to give pmethoxyphenylacetaldehyde semicarbazone, m. 180-1.5°. 1,4-Bis(pmethoxyphenyl)butane (VI), m. 77-8°, was prepared by refluxing 25 hrs. III with Zn-Hq, HCl, PhMe, and H2O, with occasional addition of HCl. Treatment of I with Na-Hg furnished 2 end products, one Et2O-insol. 1,4-bis(pmethoxyphenyl)-2-butanone (VII), m. 87-7.5, and the other Et20-soluble 1,4bis(p-methoxyphenyl)-1,3-butadiene (VIII), m. 229-31°. VII is also prepared by methylation of 1,4-bis(p-hydroxyphenyl)-2-butanone (IX), m. 185.5-6.5°, with Me2SO4. Di-Ac derivative of IX, m. 86.5-8°, was prepared by heating IX with Ac20 and AcONa. Di-Bz derivative of IX, m. 153-4°, was prepared by treatment with BzCl; 2,4-dinitrophenylhydrazone was prepared by heating VII with dinitrophenylhydrazine and H3PO4, yellow needles, m. $144-5^{\circ}$. A synthesis of VII in 4 steps is given: p-MeOC6H4CH2CN is condensed with PhCH2COCH2CN to form p-methoxyphenylacetoacetonitrile, this converted to pmethoxyphenylacetone, and the latter condensed with anisaldehyde, forming 1,4bis(p-methoxyphenyl)-3-buten-2-one, which is then bydrogenated in Me2CO in the presence of Pd black giving VIII, m. 86-7°. Bis(p-methoxybenzyl)glycolic acid (X), m. 166.5-67, was prepared from III by suspending it in KOH solution, warming to $60-70^{\circ}$ with agitation, and acidifying; the Me ester of X m. 114-15°. 1,3-Bis(p-methoxyphenyl)-2-propanone (XI), m. 85-5.5°, was made by treating AcOH solution of X with (AcO) 4Pb, warming to sep. CO2, and adding H2O to sep. the product; the oxime m. 102.5-103°. Reduction of XI by the Clemmensen procedure furnished the reduced product, m. $44-5^{\circ}$. The osazone of III was prepared by condensation of p-MeOC6H4CH2CO2Et in the presence of a fine suspension of Na in PhMe and Et2O and treatment with phenylhydrazine. Catalytic bydrogenation of VIII was carried out in semimicro apparatus using Pt from PtO, furnishing 1,4-bis(p-methoxyphenyl)butane, m. 77-8°. The presence of two isonitrile groups in I was proven quantitatively by warming an alc. solution with HgCl2 and titrating the reduced HgCl gravimetrically; in another type of determination, (CO2H)2 was treated with I and the evolved CO measured. I dimethyl ether imino ether (XII) was prepared in yellow shombs, m. $121-2^{\circ}$; the di-imino ether was formed at the same time in colorless needles, m. 193-4°. 1,4-Bis(p-methoxyphenyl)-2,3-bis(formamido)-1,3-butadiene (XIII), m. 235-6°, was prepared from I di-Me ether by boiling with glacial AcOH and adding H2O. 1,4-Bis(p-hydroxypheny1)2,3-bis(formamido)-1,3butadiene (XIV), pale yellow needles, decomposing above 200°, was prepared by warming VIII in glacial AcOH. 1,4-Bis(p-methoxyphenyl)-2,3bis(methylformamido)-1,3-butadiene (XV), m. 165-6, was prepared by treating XIV with Me2SO4 and NaOH. 1,4-Bis(p-hydroxyphenyl)-2,3bis(dimethylhydroxyacetamido) - 1,3-butadiene (XVI), m. 271-2° (decomposition), was prepared by treating I in Me2CO solution with 2N H2SO4. 1,4-Bis(p-methoxypheny1)-2,3- bis[(dimethylhydroxyacety1)methylamino]-1,3butadiene, m. 240° (decomposition), was prepared by dissolving XVI in NaOH and treating with Me2SO4 at 100° 15 min. Imidazole-4,5-dicarboxylic acid (XVII), decompose above 300° was prepared by refluxing I, I dimethyl ether, XIII, or

XIV with H2O2. The following salts of XVII were prepared (m.p. given): diethylamine (m. 180°), imidazole (m. 244°), acid K salt (m. 280°), acid Na salt, nitrate (118°), and picrate (211°). From the mother liquor remaining from the preparation of XVII, 12-15% (of theory) anisic acid was isolated, m. 184°. By hydrogenation of I with Raney Ni, 2,3-bis(p-hydroxybenzyl)pyrazine (XVIII), m.~203-4°, was obtained (diacetate, m.~91-2°, dibenzoate, m.~122-3°). 2,3-Bis(p-methoxybenzyl)pyrazine (XIX), m. 57-8°, was prepared by treating XVIII in MeOH and Et20 with CH2N2. Pyrazine-2,3-dicarboxylate was formed when XVIII was heated 2 hrs. on an H2O bath with 3% KMnO4, acidified with AcOH, vacuum-dried, heated 2 hrs. with glacial AcOH in a "bombtube," and distilled after alkalinization; picrate, m. 156-7°, Rf 0.58. 2,3-Bis(pmethoxybenzoyl)pyrazine (XX), m. 164-5° was prepared from XIX by treatment with Beckmann's mixture, m. 164-5°; the bis-(2,4-dinitrophenylhydrazone), dark red crystals, m. 299-300° (decomposition); dioxime, m. 224-9°. 3,6-Bis(pmethoxyphenyl)-1,2-diazino[4,5;2,3]pyrazine, yellow plates, m. 293.5-4.5, was prepared from XX by treatment with hydrazine sulfate and NaOH. XV in CHCl3 solution was treated with BzO2H and the rate of consumption determined XV (1 mole) consumed 2.5 moles of the acid; however, the reaction velocity after the consumption of 2 moles acid was considerably greater, indicating the presence of double bond in I. 4,5-Bis(p-methoxybenzoyl)imidazole (XXI), m. 173-4°, was prepared by oxidizing I di-Me ether with CrO3; at the same time, 2 strongly acidic substances were obtained, one being anisic acid; the bis(2,4dinitrophenylhydrazone) of XXI, dark red crystals, m. 278-9°. 3,6 - Bis(p methoxyphenyl)1,2 - diazino[4,5;4,5]imidazole (XXII), m. 146-7°, was obtained from XXI by refluxing 30 min. with hydrazine hydrate. I was not synthesized, but it was felt that the structure of I was fully established, by the results of these various studies, as [HOC6H4CH:C(NC)]2. 23 references.

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     89-01-0P, 2,3-Pyrazinedicarboxylic acid
                                              100-09-4P, p-Anisic acid
     104-01-8P, Acetic acid, (p-methoxyphenyl) - 122-84-9P, 2-Propanone,
                          4741-73-5P, Propane, 1,3-bis(p-methoxyphenyl)-
     (p-methoxyphenyl)-
     5701-83-7P, Butane, 1,4-bis(p-methoxyphenyl)-
                                                     29903-09-1P, 2-Propanone,
     1,3-bis(p-methoxyphenyl)-
                                 38565-14-9P, Formamide,
     N, N'-[bis(p-hydroxybenzylidene)ethylene]bis-
                                                    43212-67-5P, 1,3-Butadiene,
     1,4-bis(p-methoxyphenyl)- 51622-85-6P, Acetaldehyde, (p-methoxyphenyl)-,
     semicarbazone 65816-19-5P, Pyrazine, picrate 74026-69-0P, 2-Butanone,
     3-hydroxy-1,4-bis(p-methoxyphenyl)- 101110-57-0P, 2,3-Butanedione,
     1,4-bis(p-hydroxyphenyl)- 101585-09-5P, 2-Propanone,
     1,3-bis(p-methoxyphenyl)-, oxime
                                       101744-01-8P, Lactic acid,
     2-p-methoxybenzyl-3-(p-methoxyphenyl)-
                                             101869-37-8P, Imidazole,
     4,5-di-p-anisoyl- 102162-84-5P, Lactic acid,
     2-p-methoxybenzyl-3-(p-methoxyphenyl)-, methyl ester 102316-70-1P,
     Pyrazine, 2,3-bis(p-methoxybenzyl)-
                                          102590-83-0P, p-Cresol,
     \alpha-(2-p-methoxybenzyl-3-quinoxalinyl)-
                                             102704-38-1P, p-Cresol,
     \alpha, \alpha'-2, 3-quinoxalinediyldi- 102748-31-2P, 2-Butanone,
     1,4-bis(p-methoxyphenyl)-, (2,4-dinitrophenyl)hydrazone
     Quinoxaline, 2,3-bis(p-methoxybenzyl)- 103271-07-4P, Imidazole,
     4,5-di-p-anisoyl-, bis[(2,4-dinitrophenyl)hydrazone] 103986-01-2P,
     Acetoacetonitrile, 4-(p-methoxyphenyl)- 107627-09-8P, 2-Butanone,
     1,4-\text{bis}(p-\text{methoxypheny1}) - 108370-05-4P, 1H-\text{Imidazo}[4,5-d]pyridazine,
     4,7-bis(p-methoxyphenyl)- 109398-70-1P, 2,3-Butanediol,
     1,4-bis(p-methoxyphenyl)- 110435-88-6P, 3-Buten-2-one,
                               111240-82-5P, Pyrazino[2,3-d]pyridazine,
     1,4-bis(p-methoxyphenyl)-
     5,8-bis(p-methoxyphenyl) - 112271-44-0P, Formamide,
     N, N'-[bis(p-methoxybenzylidene)ethylene]bis-
                                                    113090-73-6P, Formimidic
     acid, N-(\beta-isocyano-p-methoxy-\alpha-p-methoxybenzylidenecinnamyl)-,
     methyl ester 114696-86-5P, 2,3-Butanedione, 1,4-bis(p-hydroxyphenyl)-,
     bis(phenylhydrazone) 118871-24-2P, Formimidic acid,
     N, N'-[bis(p-methoxybenzylidene)ethylene]di-, dimethyl ester
     118871-26-4P, \ \ Formamide, \ \ N, N'-[bis(p-methoxybenzylidene)ethylene]bis[N-methoxybenzylidene]
             122360-58-1P, Lactamide,
     methyl-
     N, N'-[bis(p-methoxybenzylidene)-ethylene]bis[N, 2-dimethyl- 124131-77-7P,
     Lactamide, N,N'-[bis(p-hydroxybenzylidene)-ethylene]bis[2-methyl-
     131252-64-7P, 2-Butanone, 1,4-bis(p-hydroxyphenyl)- 856638-96-5P,
     Imidazole, compound with 4,5-imidazoledicarboxylic acid
     RL: PREP (Preparation)
        (preparation of)
L9
     ANSWER 15 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                         1957:5337 CAPLUS Full-text
DOCUMENT NUMBER:
                         51:5337
ORIGINAL REFERENCE NO.: 51:1084e-i,1085a-d
                         Synthesis and reactions of guaiacylglycerol
TITLE:
                         Stumpf, Walter; Rumpf, Gunther
AUTHOR(S):
CORPORATE SOURCE:
                         Univ. Heidelberg, Germany
SOURCE:
                         Annalen der Chemie, Justus Liebigs (1956),
                         599, 51-60
                         CODEN: 9X224Y
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DOCUMENT TYPE: Journal LANGUAGE: Unavailable

OTHER SOURCE(S): CASREACT 51:5337

AΒ 3,4-MeO(PhCH2O)C6H3CHO (138 g.) in 230 cc. AcOMe, after standing overnight with 17.1 g. Na wire and 2 cc. MeOH was triturated carefully with 230 cc. AcOMe, kept another 48 hrs. at room temperature, refluxed 1 hr. with 230 cc. absolute Et20, and shaken with 740 cc. H2SO4. The organic phase washed with aqueous NaHCO3 and H2O, dried and evaporated gave 95.5% crude and 83% pure 3,4-MeO(PhCH2O)C6H3CH:CHCO2Me (I), m. 98-9° (from MeOH or PrOH). To 7.16 g. I in 24 cc. CHCl2 at 0°-5° were added dropwise (over a 1.5-hr. period) 3.84 g. Br; the mixture after 1 hr. at 0° was evaporated giving the crude di-Br derivative (II) of I (not weighed or analyzed), 22.9 g. of which were added to 15 g. dry AcOK in 150 cc. AcOH and 50 cc. Ac2O, heated 9 hrs. on a steam bath, then boiled 8 hrs., filtered, and concentrated in vacuo to incipient crystallization, treated with H2O and extracted with ET2O; the washed and dried extract evaporated in vacuo gave 19 g. sirup, a small sample of which, triturated with MeOH, gave seed crystals of (III), C22H24O8, m. sharply 112.5-13.5° (after 2 crystns. from MeOH). The main portion of the sirup, inoculated with III, crystallized gradually giving 8.25 g. of what was probably a mixture of isomeric forms of 3,4-MeO(PhCH2O)C6H3CH(OAc)CH(OAc)CO2Me (IV), m. poorly $90-100^{\circ}$ (even after repeated crystallization from MeOH). In another experiment in which 169 g. crude II was heated 15 hrs. at 100° with 110 g. AcOK, 800 cc. AcOH, and 400 cc. Ac2O, a red sirup was formed, which, with III, gave 52.45 g. IV, leaflets, m. poorly $89-98^{\circ}$, the mother liquors from which gave I. A series of fully described attempts were made to fractionate IV into its component (racemic) isomers, but although 3 fractions were obtained, m., resp., $85.5-87^{\circ}$, $86-91^{\circ}$, and $88-91^{\circ}$, none of these was homogeneous. IV (m. 89-98°)(12.5 g.) in 50 cc. dry AcOMe was hydrogenated with 0.75 g. 2% pd-BaSO4. After 1.5 hrs. 745 cc. H had been taken up. The filtered, evaporated solution gave 3,4-MeO(HO) C6H3CH(OAc) CH(OAc) CO2Me (V), viscous, uncrystallizable pale yellow sirup. V (8.34 g.) in 100 cc. absolute Et20 was added dropwise to 5.6 g. LiAlH4 in 200 cc. Et20, and after 5 hrs. at room temperature was refluxed 2 hrs., cooled to 0° in a stream of CO2 and treated dropwise with H2O, shaken with H2O saturated with CO2, the Et2O layer separated and the aqueous phase extracted continuously for 7 days with peroxide-free Et2O in a Perforator, using fresh Et2O after 28 hrs. (when a sirup separated from the Et2O-phase). The various combined Et2O exts. evaporated in vacuo gave 2.7 g. crude resinous guaiacylglycerol (VI), which was boiled briefly with 200 cc. H2O, filtered and reextd. twice with Et2O. The aqueous phase (in which VI is very soluble) was evaporated to dryness in vacuo under N, giving 1.98 g. (36.3%) purified VI, C10H14O5, yellow sirup (after drying 14 hrs. at 34° in vacuo and 2 days at 20° over P2O5). VI is difficultly soluble in Et20 and C6H6 and could not be crystallized IV (m. 87-93°) (21.2 g.) in 750 cc. absolute Et20 was stirred into a mixture of 10 g. LiAlH4 and 200 cc. Et20, and treated as in the case of V. The resultant aqueous solution was filtered and extracted 40 hrs. with Et2O; the Et2O layer yielded 2.15 g. PhCH2 derivative (VII) of VI, m. 68-74° (from C6H6), m. 99-100.5° (from MeOH by addition of Et20 and petr. ether to incipient cloudiness, or from AcOEt). Even this purified VII may be a mixture of racemic isomers. VI (1.81 g.) and Na2S2O5 in 85 cc. H2O was heated and shaken 19 hrs. at 135° in a sealed tube, then freed from SO2 and extracted 40 hrs. with Et20 giving 0.15 g. impure 4,3-HO(MeO)C6H3CH(SO3H)CH(OH)CH2OH, 81% of which was soluble in cold H2O forming a pale pink Ba salt (VIII) (containing 32.68% C and 21.3% Ba; calculated 34.71 and 19.85%, resp.). Oxidations with NaIO4 were carried out with various phenolic compds. or their derivs., and results are given in terms of moles NaIO4 consumed per mole of compound within a specific time period. No NaIO4 was consumed by veratrole within 17.5 hrs. The consumption of NaIO4 by PhOH, vanillin, and p-cresol was very slight (0.14-0.33 mole within 21.5-24 hrs.). V, VI, quaiacol, cresol, quaiacylethylcarbinol, VIII, 1,4-C6H4(OH)2, catechol, and pyrogallol all consumed appreciable amts. of NaIO4 within

- relatively short periods. On oxidation all guaiacyl compds. gave red solns., the color being ascribed to quinoid oxidation products. Oxidation data are discussed at length.
- TI Synthesis and reactions of guaiacylglycerol
- SO Annalen der Chemie, Justus Liebigs (1956), 599, 51-60 CODEN: 9X224Y
- AΒ 3,4-MeO(PhCH2O)C6H3CHO (138 g.) in 230 cc. AcOMe, after standing overnight with 17.1 q. Na wire and 2 cc. MeOH was triturated carefully with 230 cc. AcOMe, kept another 48 hrs. at room temperature, refluxed 1 hr. with 230 cc. absolute Et20, and shaken with 740 cc. H2SO4. The organic phase washed with aqueous NaHCO3 and H2O, dried and evaporated gave 95.5% crude and 83% pure 3,4-MeO(PhCH2O)C6H3CH:CHCO2Me (I), m. 98-9° (from MeOH or PrOH). To 7.16 g. I in 24 cc. CHCl2 at 0°-5° were added dropwise (over a 1.5-hr. period) 3.84 g. Br; the mixture after 1 hr. at 0° was evaporated giving the crude di-Br derivative (II) of I (not weighed or analyzed), 22.9 g. of which were added to 15 q. dry AcOK in 150 cc. AcOH and 50 cc. Ac2O, heated 9 hrs. on a steam bath, then boiled 8 hrs., filtered, and concentrated in vacuo to incipient crystallization, treated with H2O and extracted with ET2O; the washed and dried extract evaporated in vacuo gave 19 g. sirup, a small sample of which, triturated with MeOH, gave seed crystals of (III), C22H24O8, m. sharply 112.5-13.5° (after 2 crystns. from MeOH). The main portion of the sirup, inoculated with III, crystallized gradually giving 8.25 g. of what was probably a mixture of isomeric forms of 3,4-MeO(PhCH2O)C6H3CH(OAc)CH(OAc)CO2Me (IV), m. poorly 90-100° (even after repeated crystallization from MeOH). In another experiment in which 169 g. crude II was heated 15 hrs. at 100° with 110 g. AcOK, 800 cc. AcOH, and 400 cc. Ac2O, a red sirup was formed, which, with III, gave 52.45 g. IV, leaflets, m. poorly 89-98°, the mother liquors from which gave I. A series of fully described attempts were made to fractionate IV into its component (racemic) isomers, but although 3 fractions were obtained, m., resp., $85.5-87^{\circ}$, $86-91^{\circ}$, and $88-91^{\circ}$, none of these was homogeneous. IV (m. $89-98^{\circ}$)(12.5 g.) in 50 cc. dry AcOMe was hydrogenated with 0.75 g. 2% pd-BaSO4. After 1.5 hrs. 745 cc. H had been taken up. The filtered, evaporated solution gave 3,4-MeO(HO) C6H3CH(OAc) CH(OAc) CO2Me (V), viscous, uncrystallizable pale yellow sirup. V (8.34 g.) in 100 cc. absolute Et20 was added dropwise to 5.6 g. LiAlH4 in 200 cc. Et2O, and after 5 hrs. at room temperature was refluxed 2 hrs., cooled to 0° in a stream of CO2 and treated dropwise with H2O, shaken with H2O saturated with CO2, the Et2O layer separated and the aqueous phase extracted continuously for 7 days with peroxide-free Et20 in a Perforator, using fresh Et20 after 28 hrs. (when a sirup separated from the Et20-phase). The various combined Et20 exts. evaporated in vacuo gave 2.7 g. crude resinous guaiacylglycerol (VI), which was boiled briefly with 200 cc. H2O, filtered and reextd. twice with Et2O. The aqueous phase (in which VI is very soluble) was evaporated to dryness in vacuo under N, giving 1.98 g. (36.3%) purified VI, C10H14O5, yellow sirup (after drying 14 hrs. at 34° in vacuo and 2 days at 20° over P2O5). VI is difficultly soluble in Et2O and C6H6 and could not be crystallized IV (m. 87-93°) (21.2 g.) in 750 cc. absolute Et20 was stirred into a mixture of 10 g. LiAlH4 and 200 cc. Et20, and treated as in the case of V. The resultant aqueous solution was filtered and extracted 40 hrs. with Et20; the Et20 layer yielded 2.15 q. PhCH2 derivative (VII) of VI, m. 68-74° (from C6H6), m. 99-100.5° (from MeOH by addition of Et2O and petr. ether to incipient cloudiness, or from AcOEt). Even this purified VII may be a mixture of racemic isomers. VI (1.81 g.) and Na2S2O5 in 85 cc. H2O was heated and shaken 19 hrs. at 135° in a sealed tube, then freed from SO2 and extracted 40 hrs. with Et20 giving 0.15 g. impure 4,3-HO(MeO)C6H3CH(SO3H)CH(OH)CH2OH, 81% of which was soluble in cold H2O forming a pale pink Ba salt (VIII) (containing 32.68% C and 21.3% Ba; calculated 34.71 and 19.85%, resp.). Oxidations with NaIO4 were carried out with various phenolic compds. or their derivs., and results are given in terms of moles NaIO4 consumed per mole of compound within a specific time period. No NaIO4 was consumed by veratrole within 17.5 hrs. The consumption of NaIO4 by

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1T 10548-93-3P, α -Toluenesulfonic acid, α -1,2-dihydroxyethyl-4-hydroxy-3-methoxy- 66266-31-7P, α -Toluenesulfonic acid, α -1,2-dihydroxyethyl-4-hydroxy-3-methoxy-, barium salt 93878-20-7P, Cinnamic acid, 4-(benzyloxy)-3-methoxy-, methyl ester 101721-98-6P, Hydrocinnamic acid, 4-(benzyloxy)- α , β -dibromo-3-methoxy-, methyl ester 109476-22-4P, Glyceric acid, 3-(4-hydroxy-3-methoxyphenyl)-, methyl ester, 2,3-diacetate 856945-37-4P, Glyceric acid, 3-[4-(benzyloxy)-3-methoxyphenyl]-, diacetates 856945-42-1P, Glyceric acid, 3-[4-(benzyloxy)-3-methoxyphenyl]-, Me esters RL: PREP (Preparation) (preparation of)

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L9 ANSWER 16 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1955:84086 CAPLUS Full-text

DOCUMENT NUMBER: 49:84086

ORIGINAL REFERENCE NO.: 49:15798a-i,15799a-c

TITLE: Constituents of Cortex piscidiae erythrinae. II.

Synthesis of O-methylpiscidic acid

AUTHOR(S): Buckle, A. L. J.; McGookin, Alexander; Robertson,

Alexander

CORPORATE SOURCE: Univ. Liverpool, UK

SOURCE: Journal of the Chemical Society (1954)

3981-6

CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

AΒ cf. C.A. 42, 4981f. Piscidic acid was shown previouslyto be (+)-p-HOC6H4CH2C(OH)(CO2H)CH(OH)CO2H (loc. cit.) and now has been confirmed by the synthesis of p-methoxybenzyltartaric acid, identical with natural O-Me piscidic acid. Several routes for the synthesis of tartaric acids of this type were examined Et α -phenylacetoacetate, b0.8 110° (2,4dinitrophenylhydrazone, m. 94-5°), prepared by the method of Attwood, et al. (C.A. 17, 3183), with Pb(OAc)4 in HOAc, gave Et α -acetoxy- α phenylacetoacetate, b0.2 128-30°. Similarly, from Et γ -phenylacetoacetate there resulted Et α -acetoxy- γ -phenylacetoacetate (I), b0.5 143-5°. I was obtained also from phenylacetyl bromide, Et diazoacetate, and HOAc. A mixture of I, HCN, and NaOH after 12 h. was diluted with EtOH, saturated with HCl, refluxed 4 h. and filtered to remove NH4Cl. The residue after evaporation of the filtrate gave Et benzyltartrate as a mixture which was separated manually into equal amts. of racemate A, m. 174-5° (diamide, 204-6°), and racemate B, m. $194-5^{\circ}$ (diamide, m. $185-6^{\circ}$). Reduction of Et phenyloxaloacetate with moist Al amalgam gave Et β -phenylmalate, b0.04 131°, which after hydrolysis with KOH and purification from EtOAc gave β -phenylmalic acid, m. 150-60°. Fractional crystallization from EtOAc gave the racemic isomeride A, m. 172°, and from EtOAc-light petroleum (b.p. $60-80^{\circ}$) the racemic isomeride B, m. 162° . Dehydration with Ac20 of a mixture of the isomerides gave phenylmaleic anhydride, m. 120°, which on treatment with alkali gave phenylmaleic acid (II), m. $90-2^{\circ}$. II with pyridine and 0s04 in Et20 gave phenyltartaric acid, m. $173-4^{\circ}$. Oxidation of citraconic acid with NaClO3 and OsO4 gave C-

methyltartaric acid, m. 144-5° (m. 146°, given by Schmidt and Perkow, C.A. 45, 2412h); Me ester, m. $99-100^{\circ}$; diamide, m. $152-3^{\circ}$. The condensation of NaOEt and Et oxalate with Et β -p-methoxyphenylpropionate, gave Et α -ethoxalyl- β -pmethoxyphenylpropionate. After reduction with moist amalgam, there was isolated Et β -hydroxy- α -p-methoxybenzylsuccinate, b0.1. 75-7°. Hydrolysis with KOH gave a mixture of acids, m. about 130°. Fractional crystallization from EtOAc-light petroleum (b.p. $60-80^{\circ}$) gave racemate A, m. $136-7^{\circ}$, and racemate B, m. 125-6°. A mixture of these racemates with Ac2O gave pmethoxybenzylidenesuccinic anhydride (III), m. 160°, which on boiling with H20 gave p-methoxybenzylidenesuccinic acid (IV), m. and mixed m.p. 194-5° (decomposition). IV was obtained also from the condensation of anisaldehyde, Et succinate, and NaOEt. IV with Ac2O gave III, which by the boiling MeOH-H2SO4 method gave Me p-methoxybenzylidenesuccinate (V), b0.5 165°. Hydrogenation with PdCl2 catalyst of IV gave p-methoxybenzylsuccinic acid (VI) m. 98-101°, and of V gave Me p-methoxybenzylsuccinate, bl 156°, m. 35-7°. Distillation of VI at 180°/0.5 mm. gave the anhydride, m. 91-2°. From pmethoxybenzyl alc. with PCl3 in Et20 there was obtained p-methoxybenzyl chloride as an unstable oil, b25 125-7°, which on condensation with Et sodiomalonate gave Et p-methoxybenzylmalonate (VII), b0.5 145°. There was isolated from BrCH2CO2Et and VII Et α -ethoxycarbonyl- α -pmethoxybenzylsuccinate, b0.2 166-9 $^{\circ}$ which on heating with EtOH-KOH gave α $carboxy-\alpha-p-methoxybenzylsuccinic acid, m. 157-9° (decomposition). When this$ was heated at $160^{\circ}/25$ mm. for 15 min., there was obtained VI. A stirred mixture of N-bromosuccinimide, p-methoxybenzylsuccinic anhydride, benzoyl peroxide, and CC14 or CS2 as solvent, after refluxing for 12 h., evaporating the filtered mixture and extracting with EtOAc gave III. III was heated until molten, then rapidly poured on to a cold surface, the solid pulverized and refluxed with CS2, collected and washed with more solvent and the process repeated 22 times. Evaporation of the combined CS2 exts. left an orange semisolid which was extracted with Et20. The residue left on evaporation of Et20 was extracted with light petroleum (b.p. 40-60°) and on cooling, deposited p-methoxybenzylmaleic anhydride (VIII), m. 64-5°, which on recrystn. from CHCl3-light petroleum (b.p. $60-80^{\circ}$), m. $65-6^{\circ}$. VIII reverted to III on melting. Hydrolysis of VIII with H2O gave p-methoxybenzylmaleic acid (IX), m. 120° (sintered at 117°). Addition of pyridine and OsO4 to IX in Et2O and the mixture kept in a closed vessel for 3 days resulted in a brown precipitate, which after collection was treated with aqueous KOH, the solution extracted with Et20, acidified, evaporated, the residue extracted with Et20 in a Soxhlet apparatus 9 h. and the extract evaporated to obtain p-methoxybenzyltartaric acid (X), m. 205-7° (decomposition); brucine salt, $[\alpha]23.5D$ -14.39° ± 0.6° (c 2.96, 50% EtOH). Resolution of X with brucine gave p-O-methylpiscidic acid, $[\alpha]$ 23D 44.01° ± 5.0° (c 1.262, H2O), m. 169-70° (mixed m.p. with X, 173-6°); cinchonine salt, $[\alpha]17D$ 139.6° (c 6.1, EtOH); brucine salt, $[\alpha]24D$ -13.03° (c 2.131, 50% EtOH); Me ester, $[\alpha]$ 18D 78.16° (c 1.54, EtOH). The following derivs. of piscidic acid were cited: Me ester, $[\alpha]23D$ 41.52° (c 1.325, H20); Et ester, $[\alpha]17.5D$ 59.70° (c 1.551, EtOH); di-Me ester, $[\alpha]19D$ 23.71° (c 6.367, EtOH); Me p-O-benzylpiscidate, [α]19D 48.73° (c 1.786, EtOH); cinchonine salt, $[\alpha]$ 21D 146.2° (c 0.424, EtOH); (+)-Nmethylphenylisopropylamine salt, m. 179°, $[\alpha]24D$ 12.73° (c 2.09, H20). Reduction of p-methoxyphenylpyruvic acid in aqueous NaOH with 2% Na-Hg gave pmethoxyphenyllactic acid, m. 88°; Me ester (XI), b0.1 135°. Methylation of XI with Ag20 in MeI gave the Me ether of Me p-methoxyphenyllactate, b0.5 120°.

TI Constituents of Cortex piscidiae erythrinae. II. Synthesis of O-methylpiscidic acid

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    824-94-2P, Anisole, p-(chloromethyl) 886-30-6P, Succinic anhydride,
ΙT
    p-methoxybenzyl- 889-10-1P, Succinic acid, p-methoxybenzylidene-
    956-41-2P, Succinic acid, p-methoxybenzyl- 5413-05-8P, Acetoacetic acid,
    2-phenyl-, ethyl ester 6335-37-1P, Malonic acid, p-methoxybenzyl-,
    diethyl ester 10436-75-6P, p-Toluidine, N-isopropyl-, piscidate
    15542-71-9P, 1,2,2-Propanetricarboxylic acid, 3-(p-methoxyphenyl)-,
    triethyl ester 15853-34-6P, Tartaric acid, methyl-
                                                           28030-15-1P, Lactic
    acid, 3-(p-methoxyphenyl)- 42151-36-0P, Succinic acid,
    p-methoxybenzylidene-, dimethyl ester
                                            46727-01-9P, Succinic anhydride,
    p-methoxybenzylidene- 55301-58-1P, Lactic acid,
    3-(p-methoxyphenyl)-, methyl ester 76595-36-3P, Acetoacetic acid,
    2-hydroxy-4-phenyl-, ethyl ester, acetate 184242-36-2P, Oxalacetic acid,
    p-methoxybenzyl-, diethyl ester 474317-22-1P, Malic acid, 3-phenyl-,
                  792942-87-1P, Succinic acid, p-methoxybenzyl-, dimethyl
    di-Et ester
             845886-49-9P, Tartaric acid, methyl-, dimethyl ester
    855644-88-1P, Maleic anhydride, p-methoxybenzyl-
                                                        857232-37-2P.
    1,2,2-Propanetricarboxylic acid, 3-(p-methoxyphenyl)- 857559-71-8P,
    Acetoacetic acid, 2-hydroxy-2-phenyl-, ethyl ester, acetate
    858215-84-6P, Hydrocinnamic acid, p,\alpha-dimethoxy-, methyl
            860373-79-1P, Malic acid, 3-p-methoxybenzyl-, di-Et ester
    861067-75-6P, Acetoacetic acid, 2-phenyl-, ethyl ester,
    2,4-dinitrophenylhydrazone 874506-60-2P, Tartramide, 2-methyl-
    874506-62-4P, Tartaric acid, [p-(benzyloxy)benzyl]-, dimethyl
            883310-90-5P, Maleic acid, p-methoxybenzyl- 907575-67-1P,
    Cinchonine, compound with piscidic acid 907575-84-2P, Cinchonine, compound
    with p-O-methylpiscidic acid
    RL: PREP (Preparation)
        (preparation of)
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ACCESSION NUMBER: 1955:19917 CAPLUS Full-text
DOCUMENT NUMBER: 49:19917

ORIGINAL REFERENCE NO.: 49:3885i,3886a-i,3887a-e

TITLE: Synthetic experiments connected with lignin Freudenberg, Karl; Muller, Heinz G.

CORPORATE SOURCE: Univ. Heidelberg, Germany

SOURCE: Annalen der Chemie, Justus Liebigs (1953), 584, 40-53

CODEN: 9X224Y

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

OTHER SOURCE(S): CASREACT 49:19917

The following model substances are in part related to the dimers obtained by F. in expts. with coniferyl alc. (C.A. 47, 12296g), which are considered "secondary building stones" in lignin formation. Veratraldehyde condensed with CH2(CO2H)2 gave quant. yields of 3,4-(MeO)2C6H3CH:CHCO2H, whose Et ester (obtained in 94% yield) with Br in CHCl3 in artificial light yielded the dibromide, m. 110°; this was refluxed 7-8 hrs. with 3 moles KOH in alc., cooled, filtered, neutralized gradually (at about 0°) with concentrated HCl, refiltered, and concentrated in vacuo. Any salts that had been filtered, combined with those separating on concentration, were dissolved in H2O and acidified with 20% H2SO4, giving 20-30% 3,4-(MeO)2C6H3C.tplbond.CCO 2H (I), m. 156° (Fulton and Robinson, J. Chemical Society 1903, 1463). Carefully purified 4,3-Me(MeO)C6H3OH (13.8 g.) and 50 cc. MeOH containing 2.3 g. Na, evaporated in vacuo, heated 5 hrs. at 100° with 22 g. I Me ester, 50 cc. PhMe, and 13.8 g. creosol, allowed to stand 12 hrs. at room temperature, extracted with Et20, shaken repeatedly with aqueous H2SO4, and the excess creosol extracted from the Et20 with aqueous NaOH, followed by washing, drying, evaporation, and fractionation, gave 17 g. Me β -(3-methoxy-4-methylphenoxy)-3,4-dimethoxycinnamate (II), prisms, m. 107-8°. At -70 to -80°, 5 g. crude II in 100 cc. dry Et20 with 0.3 q. LiAlH4 in 26 cc. Et20 gave a precipitate which, when decomposed with H2SO4, yielded a mixture of creosol, dimethoxycinnamyl alc. (III), and α -(2-methoxy-4-methylphenoxy)-3,4dimethoxycinnamyl alcohol (IV). III and IV could not be separated by distillation, or by adsorption in C6H6 on Al2O3 but the separation was effected on a paper chromatogram with C6H6 (Rf of III and IV being 0 and 0.9, resp.). IV was noncryst., but gave a red, crystalline p-PhN2C6H4CO derivative, m. 118-19°; and a crystalline phenyl-urethan, m. 134-5°. The phenylurethan of III m. 107-8°. PhOCH2CO2Me (22 g.) and 14 g. BzH reacted vigorously with 3 g. Na wire and 40 cc. dry Et20. After 12 hrs. 8.2 g. glacial AcOH, 60 cc. H2O, and 20 cc. Et2O were added successively, giving 38% PhCH(OH)CH(OPh)CO2Na (V), the Et2O and alc. washings from which, when concentrated and esterified, yielded 48% PhCH:C(OPh)CO2Me (VI), b11 210°, m. $60-1^{\circ}$. The free acid from V, oil (not characterized) gave the Me ester (VII), m. 61° (from petr. ether); Ac derivative of VII, m. 69-70°; S-benzylthiuronium salt (corresponding to V), m. 188°. VI in Et20, under N at -70° with LiAlH4, gradually warmed to -20° with aqueous H2SO4 gave PhCH:C(OPh)CH2OH, viscous oil; phenyl urethan, m. 104°. VII, similarly reduced (at -20°) gave PhCH(OH)CH(OPh)CH2OH, b1 197°, m. 74-5°. Using Giacosa's technique [J. prakt. Chemical 19, 396(1879)] but with longer initial heating, creosol, C1CH2CO2H, and NaOH gave 67% 4,2-Me(MeO)C6H3OCH2CO2H, m. 115°; Me ester (VIII), b11 167°; amide, m. 134-5°. Veratraldehyde (15.8 g.), 20 g. VIII, and 2.2 g. powdered Na under Et20, first cooled, then heated several hrs. on a steam bath and acidified with AcOH, gave 3,4-(MeO)2C6H3CH:CRCO2Me (IX) [R in this and other compds. = 4,2-Me(MeO)C6H3O], which, reduced with LiAlH4 at -70° yielded the alc., C19H22O5 (isolated by treating the intermediate salt, under Et2O, with Dry Ice), oil, setting to a resin; 3,5-dinitrobenzoale, yellow needles, m. 158-9° (from BuOH). When 15.8 g. veratraldehyde, 20 g. VIII, 2.2 g. Na, and 50 cc. Et20 were kept at about 0° and then acidified with aqueous AcOH, the product was a mixture, b0.01 225°, of IX and 3,4-(MeO)2C6H3(OH)CHRCO2Me, m. 137° (from aqueous MeOH). To 8 g. Na (powdered under 100 cc. absolute PhMe) were added successively 25 g. abs EtOH and 50 g. vanillin, and the resulting Na derivative was filtered, triturated with and suspended in PhMe, wellcooled, and treated with freshly distilled C1CH2OMe; this kept at least 6 hrs. at room temperature, washed with 2% NaOH, and fractionated gave 41 q. methoxymethylvanillin (X), b1.5 145-7°, m. 39-40°. Freshly prepared X (9.8 g.) fused with 10.5 g. VIII, the product cooled, treated with 1.15 g. Na wire and 40 cc. Et20, allowed to stand overnight, 3.1 g. AcOH in 40 cc. H2O added, and the mixture extracted with Et20 gave 9 g. 4,3-(MeOCH2O) (MeO) C6H3CH(OH) CHRCO2Me, b0.05 175-7°. With 30 g. 14-day-old X (or with fresh X containing small amts. of vanillin), the reaction was sluggish and required heating for completion, giving as the principal product 3,4MeO(MeOCH2O)C6H3CH: CRCO2Me (XI), m. 112-13° (from aqueous MeOH). With a drop of H2SO4, AcOH, and Ac2O, 2 g. XI at 0° gave, after 1.5 hrs., 1.3 g. 3,4-MeO(AcO)C6H3CH:CRCO2Me (XII), m. 80° (from aqueous EtOH); when cooling was omitted, but the reaction continued for 8 hrs., the yield of XII was 87%. XII reduced with LiAlH4 under N at -20° , followed by a fully described extensive purification, including chromatographic fractionation on powdered cellulose, gave 3,4-MeO(HO)C6H3CH:CRCH2OH (XIII), b0.0001 140° (bath temperature), prisms, m. 90-1° (from CH2Cl2-petr. ether). Hydrogenated in MeOH with 5% Pd -BaSO4, XII gave the dihydro derivative, C21H24C7, b0.01 197°, which, reduced with LiAlH4, yielded the dihydro derivative of XIII, C18H22O5, b0.01 150°. Vanillin (10 g.), 6.25 g. CH2ClCO2H, 8.5 g. KOH, and 30 cc. H2O heated 4 hrs. at 100° and acidified with aqueous HCl gave quantitatively 2,4-MeO(OHC)C6H3OCH2CO2H, m. 188-9° [Elkan, Ber. 19, 3045(1886)], 8 g. of which with 10 g. CH2(CO2H)2 in 50 cc. pyridine containing small amts. of piperidine heated 2 hrs. at 100° yielded quantitatively 3,4-MeO(HO2CCH2O)C6H3CH:CHCO2H, m. 234° (also formed in 73% yield from ferulic acid, CH2C1CO2H and NaOH); di-Me ester (XIV), m. $104-5^{\circ}$. Veratraldehyde, (3 g.), 5 g. XIV, 0.41 g. Na powder, 20 cc. Et20, and several drops absolute MeOH, heated several hrs. and acidified with aqueous AcOH, gave 2 g. Me α -[2-methoxy-4-(β carbomethoxyvinyl)phenoxy]-3,4-dimethoxycinnamate, b0.01 260°, m. 129°. XIV and X refluxed with Na in Et2O gave, after acidification and fractionation of the Et20 extract, 31% Me 0-methoxymethy1- α -[2-methoxy-4-(β carbomethoxyvinyl)phenoxy]ferulate (XV), b0.01 275°, m. 100-1° (from MeOH). By replacing the MeOCH2 group in XV by Ac, the O-Ac analog (XVI), C24H24O9, $b0.0001\ 180^{\circ}$ (bath temperature), m. $117-18^{\circ}$, was formed. XVI (18 g.) in Et20 reduced by stepwise addition of LiAlH4 at room temperature, followed by adding moist Et20, Na2S2O4, and Dry Ice to the aqueous phase, and fractionation in high vacuum of the Et2O extract, gave about 100 mg. resinous. 3,4-MeO[3,4-MeO(HO)C6H3CH:C(CH2OH)O] C6H3CH:CHCH2OH (XVII). The tetrahydro derivative of XVI, sirup, b0.001 160° (bath temperature); the tetrahydro derivative of XVII, colorless sirup, b0.001 150° (bath temperature). Inasmuch as both acetonelignin and the dehydrogenation polymers of coniferyl alc. yielded 1.5-2% HCHO when distilled with H2SO4, a similar treatment was applied to a number of the synthetic compds. listed above. None of these gave more than faint traces of HCHO, with the single exception of PhCH(OH)CH(OPh)CH2OH, which yielded 1.3% HCHO. From this and previous studies (C.A. 42, 882a). F. and M. have indicated what types of structure, in O-containing derivs. of PhPr, are capable of giving rise to HCHO. 19 references.

- TI Synthetic experiments connected with lignin
- SO Annalen der Chemie, Justus Liebigs (1953), 584, 40-53 CODEN: 9X224Y
- The following model substances are in part related to the dimers obtained by AΒ F. in expts. with coniferyl alc. (C.A. 47, 12296g), which are considered "secondary building stones" in lignin formation. Veratraldehyde condensed with CH2(CO2H)2 gave quant. yields of 3,4-(MeO)2C6H3CH:CHCO2H, whose Et ester (obtained in 94% yield) with Br in CHCl3 in artificial light yielded the dibromide, m. 110°; this was refluxed 7-8 hrs. with 3 moles KOH in alc., cooled, filtered, neutralized gradually (at about 0°) with concentrated HCl, refiltered, and concentrated in vacuo. Any salts that had been filtered, combined with those separating on concentration, were dissolved in H2O and acidified with 20% H2SO4, giving 20-30% 3,4-(MeO)2C6H3C.tplbond.CCO 2H (I), m. 156° (Fulton and Robinson, J. Chemical Society 1903, 1463). Carefully purified 4,3-Me(MeO)C6H3OH (13.8 g.) and 50 cc. MeOH containing 2.3 g. Na, evaporated in vacuo, heated 5 hrs. at 100° with 22 g. I Me ester, 50 cc. PhMe, and 13.8 g. creosol, allowed to stand 12 hrs. at room temperature, extracted with Et20, shaken repeatedly with aqueous H2SO4, and the excess creosol extracted from the Et20 with aqueous NaOH, followed by washing, drying, evaporation, and fractionation, gave 17 g. Me β -(3-methoxy-4-methylphenoxy)-3,4-dimethoxycinnamate (II), prisms, m. 107-8°. At -70 to -80°, 5 g. crude II

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ΤТ
     1660-19-1P, Acetic acid, (4-formyl-2-methoxyphenoxy)-
                                                              2316-26-9P,
     Cinnamic acid, 3,4-dimethoxy- 5533-00-6P, Veratraldehyde,
                   6270-23-1P, Acetic acid, (2-methoxy-p-tolyloxy)-
     18523-76-7P, 2-Propen-1-ol, 3-(3,4-dimethoxyphenyl)- 20583-78-2P,
     Cinnamic acid, 3,4-dimethoxy-, ethyl ester 22511-06-4P, Propiolic acid,
     (3,4-dimethoxyphenyl) - 41247-45-4P, 2-Propen-1-ol,
     3-(4-hydroxy-3-methoxyphenyl)-2-(2-methoxy-p-tolyloxy)-
     1-Propanol, 3-(4-hydroxy-3-methoxypheny1)-2-[4-(3-hydroxypropy1)-2-
     methoxyphenoxy] - 62497-24-9P, Propiolic acid, (3,4-dimethoxyphenyl)-,
     methyl ester 70110-65-5P, 1,3-Propanediol, 2-phenoxy-1-phenyl-
     84159-61-5P, Acetic acid, (2-methoxy-p-tolyloxy)-, methyl ester
     99873-50-4P, Hydrocinnamic acid, \alpha, \beta-dibromo-3, 4-dimethoxy-,
     ethyl ester 100519-48-0P, Cinnamic acid, 4-(carboxymethoxy)-3-methoxy-
     102448-71-5P, Ferulic acid, \alpha-(2-methoxy-p-tolyloxy)-, methyl ester,
               102448-71-5P, Hydroferulic acid,
     acetate
     \alpha-(2-methoxy-p-tolyloxy)-, methyl ester, acetate 102596-16-7P,
     Cinnamic acid, 3-methoxy-4-(methoxymethoxy)-\alpha-(2-methoxy-p-tolyloxy)-
     , methyl ester
                     102749-35-9P, Cinnamic acid,
     4-hydroxy-\alpha, 4'-oxybis[3-methoxy-, dimethyl ester, acetate
     102749-35-9P, Ferulic acid, \alpha-[4-(2-carboxyviny1)-2-methoxyphenoxy]-
     , dimethyl ester, acetate 198897-12-0P, Cinnamic acid,
     4-(carboxymethoxy)-3-methoxy-, dimethyl ester 854884-24-5P, Cinnamic
     acid, \alpha-phenoxy-, methyl ester
                                     854888-06-5P, Cinnamic acid,
     3,3',4-trimethoxy-\alpha,4'-oxydi-, dimethyl ester
                                                     856179-22-1P,
     Cinnamic acid, 4-(methoxymethoxy)-\alpha, 4'-oxybis[3-methoxy-, dimethyl]
           856818-05-8P, 1-Propanol, 3-(4-hydroxy-3-methoxyphenyl)-2-(2-
                          857231-66-4P, 2-Propen-1-ol, 2-phenoxy-3-phenyl-,
     methoxy-p-tolyloxy)-
     carbanilate 857234-56-1P, 2-Propen-1-ol,
     3-(4-hydroxy-3-methoxyphenyl)-2-[4-(3-hydroxypropenyl)-2-methoxyphenoxy]-
     857946-32-8P, Acetamide, 2-(2-methoxy-p-tolyloxy)- 858207-72-4P,
     Hydracrylic acid, 3-(3,4-dimethoxyphenyl)-2-(2-methoxy-p-tolyloxy)-,
     methyl ester 858207-82-6P, Hydracrylic acid,
     3-[3-methoxy-4-(methoxymethoxy)phenyl]-2-(2-methoxy-p-tolyloxy)-, methyl
     ester 858217-33-1P, Hydrocinnamic acid,
     4-hydroxy-\alpha, 4'-oxybis[3-methoxy-, dimethyl ester, acetate
     859056-37-4P, 2-Propen-1-ol, 2-phenoxy-3-phenyl- 859056-69-2P,
     2-Propen-1-ol, 3-(3,4-dimethoxyphenyl)-2-(2-methoxy-p-tolyloxy)-,
     3,5-dinitrobenzoate 859056-72-7P, 2-Propen-1-ol,
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3-(3,4-dimethoxyphenyl)-, carbanilate 859060-89-2P, Pseudourea, 2-benzyl-2-thio-, compound with 2-phenoxy-3-phenylhydracrylic acid 860257-90-5P, Hydroferulic acid, α -[4-(2-carboxyethyl)-3-methoxyphenoxy]-, dimethyl ester, acetate 873410-58-3P, Benzoic acid, p-phenylazo-, 3,4-dimethoxy- γ -(2-methoxy-p-tolyloxy)cinnamyl ester 874009-71-9P, 2-Propen-1-ol, 3-(3,4-dimethoxyphenyl)-2-(2-methoxy-p-tolyloxy)- RL: PREP (Preparation) (preparation of)

L9 ANSWER 18 OF 18 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1953:72704 CAPLUS Full-text

DOCUMENT NUMBER: 47:72704

ORIGINAL REFERENCE NO.: 47:12323b-i,12324a-h

TITLE: Aliphatic nitro compounds. IV. Various addition and

condensation reactions of ethyl nitroacetate

AUTHOR(S): Dornow, Alfred; Frese, Albert

CORPORATE SOURCE: Tech. Hochschule, Hannover, Germany

SOURCE: Annalen der Chemie, Justus Liebigs (1952),

578, 122-36 CODEN: 9X224Y

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

OTHER SOURCE(S): CASREACT 47:72704 GΙ For diagram(s), see printed CA Issue. AΒ O2NCH2CO2Et (I) (prepared by Steinkopf's method, C.A. 18, 377) added α, β unsatd. ketones at room temperature, with Et2NH (II) or PhCH2NH3OH (III) as catalysts. In ligroine, I and PhCH: CHAc with II gave 92% [AcCH2CHPhC(NO2)(CO2Et)]NH2Et2 (IV), m. 114°. Similarly I, II, and PhCH2:CHNO2 gave the NH2Et2 salt of Et 2-aci-2,4-dinitro-3-phenylbutyrate (V), m. 98° (and traces of polynitrostyrene). V (96% was also formed from equivalent amts. of Ph(PhNH)CHCH2NO2, I, and II, in ligroine and Et2O; in the absence of Et20, polynitrostyrene is a by-product. I (3.7 g.), 2-cc. of 40% III, 4 cc. dioxane, and 4 g. PhCH:CHCO2Et, kept 3 days at 60° , then at 0° , and the mixture acidified with N HCl and extracted with Et20 gave 66% EtO2CCH(NO2)CHPhCH2CO2Et, b1 176-8°. PhCH:CHAc and I with III gave, after 14 days, a brown smear from which only 30 mg. of the PhCh2NMe2 analog of V, m. 175° , could be isolated by Et20 extraction. The residual Et20 extract treated with II gave 54% V. I, II, and (PhN:)2 yielded 82% of the adduct C20H28O4N4, m. 88°. In ligroine or petr. ether, I with a variety of Schiff bases in the presence of II yielded unstable adducts [R1HNCHRC(NO2)CO2Et]NH2Et2 (VI), in which R and R1 are usually aryl groups, given in the following table. Schiff base used, Yield of, VI %, Decomposition, °C., Formula of VI; PhN:CHPh, 99, 106, (VIa) C21H29O4N3; p-MeC6H4N:CHPh, 94, 86, (VIb) C22H31O4N3; p-EtOC6H4N:CHPh, 96, 106, (VIc) C23H33O5N3; m-O2NC6H4N:CHPh, 93, 109, (VId) C21H28O6N4; p-O2NC6H4N:CHPh, 95, 120, (VIe) C21H28O6N4; MeN:CHPh, 97, 78, (VIf) C16H27O4N3; o-O2NC6H4CH:NPh, 95, 105, (VIg) C21H27O6N4; m-O2NC6H4CH:NPh, 96, 113, (VIh) C21H28O6N4; p-O2NC6H4CH:NPh, 86, 105, (VIi) C21H28O6N4; m-O2NC6H4CH:NC6H4Me-p, Too unstable for isolation; o-HOC6H4CH:NPh, " " " "; 2,1-HOC10H6CH:NPh, " " " "; p-HOC6H4CH:NPh, " " " "; p-MeOC6H4CH:NPh, " " " "; o-ClC6H4CH:NPh, 98, 108, (VIj) C21H28O4N3Cl; p-ClC6H4CH:NPh, 99, 84, (VIk) C21H28O4N3Cl; N-(2-Pyridylmethylene)-p-phenetidine, 95, 86, (VII) C22H32O5N4; N-(3-Pyridylmethylene)-p-phenetidine, 82, 103, (VIm) C22H32O5N4; N-mathematical National Nafurfurylidene-p-phenetidine, 85, 112, (VIn) C21H31O6N3; Attempts to precipitate the VI in CHCl3 with petr. ether, or solution in alc. (or Et20) caused decomposition to the corresponding Et2NH salts (VII) of RCH[CH(NO2)CO2Et]2 (R = aryl). Thus VIa, VIb, VIc, VId, and VIe yielded resp. 94-98, 95, 92, 89, and 84% VIIa, C19H29O8N3, m. 129° (decomposition). VIg gave 82% (VIIb) C19H28O10N4, m. 126°. VIh gave 90% VIIc C19H28O10N4, m. 121°.

VIi gave 82% VIId, C19H28O10N4, m. 125° . From m-O2NC6H4CH:NC6H4Me-p was formed 95% VIIc, m. 121°; from o-HOC6H4CH:NPh, 90% VIIe, C19H29O9N3, m. 128°; from p-HOC6H4CH:NPh, 64%, VIIf, C19H29O9N3, m. 119°; from p-MeOC6H4CH:NPh, 72% VIIg, C20H3109N3, m. 95-7°. The product from 2-HOC10H6CH:NPh could not be isolated. VIj gave 97% VIIh, C19H28O8N3Cl, m. 118°; VIk, 67% VIIj; C19H28O8N3Cl, m. 125°; VII, 84% VIIk, C18H28O8N4, m. 139°; VIm, 55% VIII, C18H28O8N4, m. 148°; and VIn, 88% VIIm, C17H27O9N3, m. 125°. When compds. of type VI in alc. were heated, preferably with an excess II, the following corresponding isoxazoline oxides (VIII) [all m. (decomposition)] were formed (cf. preceding abstract): from VIa, 76-80% C17H22O5N2, m. 181°; from VIf, 78% C14H16O5N2, m. 181°; from VIg, 38% C17H21O7N3, (VIIIa), m. 189°; from VIh, 46% of the m-isomer of VIIIa, m. 181° ; from VIi, 32% of the p-isomer of VIIIa, m. 187°; from VIj, 68% C17H21O5N2Cl, m. 196°; from VIk, 70% of the p-isomer, m. 168°. Although the corresponding VI and VII from 2-HOC10H6CH:NPh could not be isolated, D. and F. claim the formation of 73% VIII from this Schiff base but give the isoxazoline oxide formula as "C23H29O5N3" m. 162° (decomposition). o-HOC6H4CH:NPh gave the isoxazoline oxide, C17H22O6N2 (incorrectly given as "C17H22O5N2"), m. 198° (decomposition). p-MeOC6H4CH:NPh gave C18H24O6N2, m. 195° (decomposition). VIIc refluxed 15 min. in absolute alc. with a large excess of II gave 42% O←N:C(CONEt2).CH(C6H4NO2).CH(CONEt2).O, m. 203° (from EtOH). Aldehydes or Schiff bases react with Et2NH2+ salts of I, giving, in the case of BzH, o-HOC6H4CH:NPh, or N-(2-pyridylmethylene)-p-phenetidine, compds. of type VII. However PhCH:NPh or its m-NO2 derivative (IX), gave directly the corresponding VIII. With the MeNH3+ salt (X) of I, a compound of type VII, C16H22O10N4, m. 130°, was obtained from IX. In all other cases X gave the following compds. of type VIII: from BzH or PhCH:NPh, 78-86% C14H16O5N2, m. 181-2° (decomposition); from p-HOC6H4CH:NPh, 62% C14H16O6N2, m. 197° (decomposition); from p-MeOC6H4CH:NPh, 57% C15H18O6N2, m. 180° (decomposition). Hydrobenzamide (3 g.) treated in ligroine at 0° with 2.6 g. I and the crystalline deposit promptly separated and washed with Et20 gave 0.7 g. NH4 salt of I, m. 124° . If the salt is not separated, it forms an oil, resolidifying in 24 hrs. and giving 2.5 g. of the di-NH4 salt of di-Et 2,4dinitro-3-phenylglutarate, C15H24O8N4, m. 95-6°. I (2 moles), 1.5 moles II, and 1,1'-benzylidenedipiperidine (XI) in Et2O-alc. (1:1) at room temperature gave 93% VIIa, m. 129°. Similarly the o-HO analog of XI gave VIIe. Cl3CCH: NC6H4Me in alc.-Et2O with I and II gave 93% [CCl3CH(OH)C(NO2)(CO2Et)](NH2Et2), m. 126° (decomposition). Cyclohexanone (5 g.), 6 g. I, and 3 cc. of 25% PhCH2NMe3OBu (XII) in BuOH gave 3.9 g. CH2.(CH2)4.C[CH(NO2)CO2Et]2, b0.8 120°. BzMe and I with XII gave 70% EtO2C.CH(NO2)CMePhCH(NO2)CO2Et, b1 150° O \leftarrow N:C(CONEt2).CH(C6H4OHp).CH(CO2Et).O in Et2O at 0° with HCl gas gave 83% O.N: C(CO2H).C(C6H4OHp):CCO2Et(XIII),m. 155° (from C6H6). Hydrogenated in the presence Pd-BaSO4 or PtO2 and MeOH, XIII gave 83% HON:C(CONEt2) CH(C6H4OH) CH(OH) CO2Et, yellow, m. 157° (decomposition); the Ph analog, m. 140° (decomposition).

- TI Aliphatic nitro compounds. IV. Various addition and condensation reactions of ethyl nitroacetate
- SO Annalen der Chemie, Justus Liebigs (1952), 578, 122-36 CODEN: 9X224Y
- AB O2NCH2CO2Et (I) (prepared by Steinkopf's method, C.A. 18, 377) added α , β -unsatd. ketones at room temperature, with Et2NH (II) or PhCH2NH3OH (III) as catalysts. In ligroine, I and PhCH:CHAc with II gave 92% [AcCH2CHPhC(NO2)(CO2Et)]NH2Et2 (IV), m. 114°. Similarly I, II, and PhCH2:CHNO2 gave the NH2Et2 salt of Et 2-aci-2,4-dinitro-3-phenylbutyrate (V), m. 98° (and traces of polynitrostyrene). V (96% was also formed from equivalent amts. of Ph(PhNH)CHCH2NO2, I, and II, in ligroine and Et2O; in the absence of Et2O, polynitrostyrene is a by-product. I (3.7 g.), 2-cc. of 40% III, 4 cc. dioxane, and 4 g. PhCH:CHCO2Et, kept 3 days at 60°, then at 0°, and the mixture acidified with N HCl and extracted with Et2O gave 66% EtO2CCH(NO2)CHPhCH2CO2Et, bl 176-8°. PhCH:CHAc and I with III gave, after 14

days, a brown smear from which only 30 mg. of the PhCh2NMe2 analog of V, m. 175°, could be isolated by Et20 extraction The residual Et20 extract treated with II gave 54% V. I, II, and (PhN:)2 yielded 82% of the adduct C20H28O4N4, m. 88° . In ligroine or petr. ether, I with a variety of Schiff bases in the presence of II yielded unstable adducts [R1HNCHRC(NO2)CO2Et]NH2Et2 (VI), in which R and R1 are usually aryl groups, given in the following table. Schiff base used, Yield of, VI %, Decomposition, °C., Formula of VI; PhN:CHPh, 99, 106, (VIa) C21H29O4N3; p-MeC6H4N:CHPh, 94, 86, (VIb) C22H31O4N3; p-EtOC6H4N:CHPh, 96, 106, (VIc) C23H33O5N3; m-O2NC6H4N:CHPh, 93, 109, (VId) C21H28O6N4; p-O2NC6H4N:CHPh, 95, 120, (VIe) C21H28O6N4; MeN:CHPh, 97, 78, (VIf) C16H27O4N3; o-O2NC6H4CH:NPh, 95, 105, (VIg) C21H27O6N4; m-O2NC6H4CH:NPh, 96, 113, (VIh) C21H28O6N4; p-O2NC6H4CH:NPh, 86, 105, (VIi) C21H28O6N4; m-O2NC6H4CH:NC6H4Me-p, Too unstable for isolation; o-HOC6H4CH:NPh, " " " "; 2,1-HOC10H6CH:NPh, " " " "; p-HOC6H4CH:NPh, " " " "; p-MeOC6H4CH:NPh, " " " "; o-ClC6H4CH:NPh, 98, 108, (VIj) C21H28O4N3Cl; p-ClC6H4CH:NPh, 99, 84, (VIk) C21H28O4N3Cl; N-(2-Pyridylmethylene)-p-phenetidine, 95, 86, (VII) C22H32O5N4; N-(3-Pyridylmethylene)-p-phenetidine, 82, 103, (VIm) C22H32O5N4; Nfurfurylidene-p-phenetidine, 85, 112, (VIn) C21H31O6N3; Attempts to precipitate the VI in CHCl3 with petr. ether, or solution in alc. (or Et20) caused decomposition to the corresponding Et2NH salts (VII) of RCH[CH(NO2)CO2Et]2 (R = aryl). Thus VIa, VIb, VIc, VId, and VIe yielded resp. 94-98, 95, 92, 89, and 84% VIIa, C19H29O8N3, m. 129° (decomposition). VIg gave 82% (VIIb) C19H28O10N4, m. 126°. VIh gave 90% VIIc C19H28O10N4, m. 121°. VIi gave 82% VIId, C19H28O10N4, m. 125°. From m-O2NC6H4CH:NC6H4Me-p was formed 95% VIIc, m. 121°; from o-HOC6H4CH:NPh, 90% VIIe, C19H29O9N3, m. 128°; from p-HOC6H4CH:NPh, 64%, VIIf, C19H29O9N3, m. 119°; from p-MeOC6H4CH:NPh, 72% VIIg, C20H31O9N3, m. 95-7°. The product from 2-HOC10H6CH:NPh could not be isolated. VIj gave 97% VIIh, C19H28O8N3Cl, m. 118°; VIk, 67% VIIj; C19H28O8N3Cl, m. 125°; VII, 84% VIIk, C18H28O8N4, m. 139°; VIm, 55% VIII, C18H28O8N4, m. 148°; and VIn, 88% VIIm, C17H27O9N3, m. 125°. When compds. of type VI in alc. were heated, preferably with an excess II, the following corresponding isoxazoline oxides (VIII) [all m. (decomposition)] were formed (cf. preceding abstract): from VIa, 76-80% C17H22O5N2, m. 181°; from VIf, 78% C14H16O5N2, m. 181°; from VIg, 38% C17H21O7N3, (VIIIa), m. 189°; from VIh, 46% of the m-isomer of VIIIa, m. 181°; from VIi, 32% of the p-isomer of VIIIa, m. 187°; from VIj, 68% C17H21O5N2Cl, m. 196°; from VIk, 70% of the p-isomer, m. 168°. Although the corresponding VI and VII from 2-HOC10H6CH:NPh could not be isolated, D. and F. claim the formation of 73% VIII from this Schiff base but give the isoxazoline oxide formula as "C23H29O5N3" m. 162° (decomposition). o-HOC6H4CH:NPh gave the isoxazoline oxide, C17H22O6N2 (incorrectly given as "C17H22O5N2"), m. 198° (decomposition). p-MeOC6H4CH:NPh gave C18H24O6N2, m. 195° (decomposition). VIIc refluxed 15 min. in absolute alc. with a large excess of II gave 42% O←N:C(CONEt2).CH(C6H4NO2).CH(CONEt2).O, m. 203° (from EtOH). Aldehydes or Schiff bases react with Et2NH2+ salts of I, giving, in the case of BzH, o-HOC6H4CH:NPh, or N-(2-pyridylmethylene)-p-phenetidine, compds. of type VII. However PhCH:NPh or its m-NO2 derivative (IX), gave directly the corresponding VIII. With the MeNH3+ salt (X) of I, a compound of type VII, C16H22O10N4, m. 130°, was obtained from IX. In all other cases X gave the following compds. of type VIII: from BzH or PhCH:NPh, 78-86% C14H16O5N2, m. 181-2° (decomposition); from p-HOC6H4CH:NPh, 62% C14H16O6N2, m. 197° (decomposition); from p-MeOC6H4CH:NPh, 57% C15H18O6N2, m. 180° (decomposition). Hydrobenzamide (3 g.) treated in ligroine at 0° with 2.6 g. I and the crystalline deposit promptly separated and washed with Et20 gave 0.7 g. NH4 salt of I, m. 124°. If the salt is not separated, it forms an oil, resolidifying in 24 hrs. and giving 2.5 g. of the di-NH4 salt of di-Et 2,4dinitro-3-phenylglutarate, C15H24O8N4, m. 95-6°. I (2 moles), 1.5 moles II, and 1,1'-benzylidenedipiperidine (XI) in Et20-alc. (1:1) at room temperature gave 93% VIIa, m. 129°. Similarly the o-HO analog of XI gave VIIe. p-Cl3CCH:NC6H4Me in alc.-Et2O with I and II gave 93% [CCl3CH(OH)C(NO2)(CO2Et)](NH2Et2), m. 126° (decomposition). Cyclohexanone (5

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CH2.(CH2)4.C[CH(NO2)CO2Et]2, b0.8 120°. BzMe and I with XII gave 70%
     EtO2C.CH(NO2)CMePhCH(NO2)CO2Et, b1 150^{\circ} O\leftarrowN:C(CONEt2).CH(C6H4OH-
     p).CH(CO2Et).O in Et20 at 0° with HCl gas gave 83% O.N: C(CO2H).C(C6H4OH-
     p):CCO2Et(XIII), m. 155° (from C6H6). Hydrogenated in the presence Pd-BaSO4 or
     PtO2 and MeOH, XIII gave 83% HON:C(CONEt2) CH(C6H4OH) CH(OH) CO2Et, yellow, m.
     157° (decomposition); the Ph analog, m. 140° (decomposition).
     521942-14-3P, 1,1-Cyclohexanediacetic acid, \alpha,\alpha'-dinitro-,
     diethyl ester 854704-42-0P, Glutaramic acid,
     N, N-diethyl-2-hydroxy-3-(p-hydroxyphenyl)-4-oxo-, ethyl ester, oxime
     854705-67-2P, Glutaric acid, 2-nitro-3-phenyl-, diethyl ester
     855601-37-5P, 2-Isoxazoline-3,5-dicarboxamide,
     N, N', N', N'-tetraethyl-4-(m-nitro-phenyl)-, 2-oxide 855601-50-2P,
     2-Isoxazoline-5-carboxylic acid, 4-(p-hydroxyphenyl)-3-methyl-carbamoyl-,
     ethyl ester, 2-oxide 855747-58-9P, 2-Isoxazoline-5-carboxylic acid,
     3-diethylcarbamoyl-4-[p-nitrophenyl]-, ethyl ester, 2-oxide
     855747-59-0P, 2-Isoxazoline-5-carboxylic acid,
     3-diethylcarbamoyl-4-[o-nitrophenyl]-, ethyl ester, 2-oxide
     855747-60-3P, 2-Isoxazoline-5-carboxylic acid,
     3-diethylcarbamoyl-4-[m-nitrophenyl]-, ethyl ester, 2-oxide
     855747-62-5P, 2-Isoxazoline-5-carboxylic acid,
     3-diethylcarbamoyl-4-(o-hydroxyphenyl)-, ethyl ester, 2-oxide
     857557-42-7P, Acetic acid, aci-nitro-, ethyl ester, NH4 derivative
     858250-71-2P, Glutaric acid, 3-methyl-2,4-dinitro-3-phenyl-, diethyl ester
     859305-07-0P, Glutaramic acid, N,N-diethyl-2-hydroxy-4-oxo-3-phenyl-,
     ethyl ester, oxime 860372-21-0P, 2-Isoxazoline-5-carboxylic acid,
     3-diethylcarbamoyl-4-(2-hydroxy-1-naphthyl)-, ethyl ester, 2-oxide
     872788-69-7P, 2-Isoxazoline-5-carboxylic acid,
                                                        872788-69-7P,
     3-methylcarbamoyl-4-phenyl-, ethyl ester, 2-oxide
     2-Isoxazoline-5-carboxylic acid, 3-methylcarbamoyl-4-phenyl-, ethyl ester,
             872788-71-1P, 2-Isoxazoline-5-carboxylic acid,
     2-oxide
     4-(p-methoxyphenyl)-3-methyl-carbamovl-, ethyl ester, 2-oxide
     872788-73-3P, 2-Isoxazoline-5-carboxylic acid,
     3-diethylcarbamoyl-4-phenyl-, ethyl ester, 2-oxide
                                                         872788-75-5P.
     2-Isoxazoline-5-carboxylic acid, 3-diethylcarbamoyl-4-(p-methoxyphenyl)-,
     ethyl ester, 2-oxide 872788-77-7P, 2-Isoxazoline-5-carboxylic acid,
     4-[p-chlorophenyl]-3-diethylcarbamoyl-, ethyl ester, 2-oxide
     874533-61-6P, 2-Isoxazoline-5-carboxylic acid,
     4-[o-chlorophenyl]-3-diethylcarbamoyl-, ethyl ester, 2-oxide
     RL: PREP (Preparation)
        (preparation of)
L12
        STRUCTURE UPLOADED
=> s 112 sss sam
SAMPLE SEARCH INITIATED 13:15:50 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED -
                                     2561 TO ITERATE
 78.1% PROCESSED
                     2000 ITERATIONS
                                                                 21 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
FULL FILE PROJECTIONS: ONLINE **COMPLETE**
                        BATCH **COMPLETE**
PROJECTED ITERATIONS:
                             48185 TO 54255
PROJECTED ANSWERS:
                               226 TO
                                           848
L13
             21 SEA SSS SAM L12
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=> d 113

q.), 6 q. I, and 3 cc. of 25% PhCH2NMe3OBu (XII) in BuOH gave 3.9 q.

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RN
    1089316-65-3 REGISTRY
ΕD
     Entered STN: 24 Dec 2008
CN
     Benzenepropanoic acid, 4-hydroxy-3-methoxy-\alpha-(2-methoxy-4-
     propylphenoxy)-\beta-oxo-\alpha-(phenylmethyl)-, ethyl ester (CA INDEX
     NAME)
MF
     C29 H32 O7
SR
     CA
/ Structure 112 in file .gra /
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
=> s 112 sss full
FULL SEARCH INITIATED 13:16:55 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 50873 TO ITERATE
100.0% PROCESSED
                    50873 ITERATIONS
                                                                 499 ANSWERS
SEARCH TIME: 00.00.01
L14
            499 SEA SSS FUL L12
=> d scan
L14 499 ANSWERS
                   REGISTRY COPYRIGHT 2009 ACS on STN
     Butanedioic acid, 2-[3-(3,4-dihydroxyphenyl)-2-hydroxy-1-oxopropoxy]-3-[[3-
     (3,4-dihydroxyphenyl)-1-oxo-2-propen-1-yl]oxy]-
     C22 H20 O13
MF
=> s 114/prep
           450 L14
       4706241 PREP/RL
L15
           224 L14/PREP
                 (L14 (L) PREP/RL)
=> s 115 and (py<2003 or ay<2003 or pry<2003)
      22983071 PY<2003
       4502933 AY<2003
       3971676 PRY<2003
L16
           142 L15 AND (PY<2003 OR AY<2003 OR PRY<2003)
=> s 116 and (asymmetri? hydrogenati?)
         93416 ASYMMETRI?
        147845 ASYM
             6 ASYMS
        147848 ASYM
                 (ASYM OR ASYMS)
        184557 ASYMMETRI?
                 (ASYMMETRI? OR ASYM)
        187623 HYDROGENATI?
          4068 ASYMMETRI? HYDROGENATI?
                 (ASYMMETRI?(W) HYDROGENATI?)
L17
             0 L16 AND (ASYMMETRI? HYDROGENATI?)
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L13 ANSWER 1 OF 21 REGISTRY COPYRIGHT 2009 ACS on STN

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=> s 116 and asymmetric? hydrogenation?
        82387 ASYMMETRIC?
        147845 ASYM
            6 ASYMS
        147848 ASYM
                (ASYM OR ASYMS)
       174519 ASYMMETRIC?
                (ASYMMETRIC? OR ASYM)
        184506 HYDROGENATION?
          4063 ASYMMETRIC? HYDROGENATION?
                 (ASYMMETRIC?(W) HYDROGENATION?)
            0 L16 AND ASYMMETRIC? HYDROGENATION?
L18
=> s 116 and 'chiral? cataly?'
       129207 'CHIRAL'
           19 'CHIRALS'
        129212 'CHIRAL'
                ('CHIRAL' OR 'CHIRALS')
            2 'CATALY'
             0 'CHIRAL? CATALY?'
                ('CHIRAL'(W)'CATALY')
            0 L16 AND 'CHIRAL? CATALY?'
L19
=> s 116 and (Ru? or Rh? or Pd? or Ir?
UNMATCHED LEFT PARENTHESIS 'AND (RU?'
The number of right parentheses in a query must be equal to the
number of left parentheses.
=> s 116 and (Ru? or Rh? or Pd? or Ir?)
       1446878 RU?
       783851 RH?
       329541 PD?
       2695423 IR?
L20
           26 L16 AND (RU? OR RH? OR PD? OR IR?)
=> d 120 ibib abs 1-5
L20 ANSWER 1 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2003:855905 CAPLUS Full-text
DOCUMENT NUMBER:
                        139:350938
TITLE:
                        Preparation of N-cinnamoyl-DOPA esters and related
                        compounds as T lymphocyte inhibitors
                        Won, Jongwha; Lee, Keunhyeung; Park, Seehyoung; Kim,
INVENTOR(S):
                        Sung-Joo; Yun, Su-Young; Kang, Mi-Ae; Hur, Yun-Gyoung;
                        Youn, Jeehee; Yun, Yungdae; Park, Doohong; Oh, Jaetaek
                        Mogam Biotechnology Research Institute, S. Korea
PATENT ASSIGNEE(S):
SOURCE:
                        PCT Int. Appl., 109 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
                  KIND
                                      APPLICATION NO. DATE
    PATENT NO.
                               DATE
    WO 2003089405
                        A1 20031030 WO 2003-KR751
                                                                20030414 <--
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KZ, LC, LK, LR, LS,
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LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL,

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PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA,
             UG, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     US 20040082664
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PRIORITY APPLN. INFO.:
                                            KR 2002-20481
                                            WO 2003-KR751
                                                              W 20030414
OTHER SOURCE(S):
                       MARPAT 139:350938
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GΙ

/ Structure 118 in file .gra /

AΒ Title compds. I [R1-R10 = H, OH, halogen, alkoxy, CHO, CO2H, NH2, CF3, NO2, \geq 1 of R1-R5 and R6-R10 = OH; X1 = O, S, NH, NMe, NEt, NHNH; X2 = CH2, CO, CS, CONH; X3 = bond, (un)substituted CH:CH, CH:CHCH:CH, CH2, CH2CH2; Y1 = H, CH2, CO, CS, alkyl, amino, 3-methyl-1,2,4-oxadiazol-5-yl, 3-benzyl-1,2,4-oxadiazol-5-yl; Y = absent, (un)substituted NH2, OH, SH; B = H, alkyl] were prepared as inhibitors of the activation of T lymphocytes by the src homol. region 2(SH2) domain of T lymphocyte (lck), useful for the treatment, prevention and/or diagnosis of graft rejection, autoimmune diseases, inflammatory diseases, etc. Thus, D-DOPA was converted to its Me ester and treated with caffeic acid to give the amide II which inhibited the binding of the lck SH2 domain with its cognate peptide < 10µM.

REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS 3 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L20 ANSWER 2 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN 2003:570940 CAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 139:133345

TITLE: Preparation of Phenyl(alkyl)carboxylic acid

derivatives and analogs and their serum glucose and/or

serum lipid lowering activity

Giannessi, Fabio; Tassoni, Emanuela; Dell'Uomo, INVENTOR(S):

Natalina; Brunetti, Tiziana; Tinti, Maria Ornella;

Arduini, Arduino; Pessotto, Pompeo

PATENT ASSIGNEE(S): Sigma-Tau Industrie Farmaceutiche Riunite S.p.A.,

Italy

SOURCE: PCT Int. Appl., 97 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Pat.ent. LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 2003059864	A2 200307	724 WO 2003-IT7	20030113 <
WO 2003059864	A3 200401	∟29	
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CO, CR, CU,	CZ, DE, DK, D	DM, DZ, EC, EE, ES, FI,	GB, GD, GE, GH,

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US 20050032787 A1 20050210 US 2004-501135 20040713 <--
RITY APPLN. INFO.:

IT 2002-RM16 A 20020115 <--
WO 2003-IT7 W 20030113
PRIORITY APPLN. INFO.:
OTHER SOURCE(S): MARPAT 139:133345
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/ Structure 119 in file .gra /

AB Title compds. I [A = CH; alkanylidene with 2-4 C atoms, etc.; Ar = mono/bicyclic (hetero)aryl; f, h = 0-1; m = 0-3; n = 0-1 and if n = 0, R1 = absent and COY is directly bound to benzene; Q, Z = NH, O, S, NHCO, etc.; Y = OH, alkoxy, amino] are prepared For instance, 3-hydroxybenzaldehyde is condensed with dimethylmalonate (HOAc, piperidine, 5 h) and the product reduced (MeOH, H2-10% Pd/C @ 50 psi, 18 h) to give II. II is capable of increasing glucose consumption in 3T3 - L1 cells to a similar extent to that achieved by rosiglitazone. I are serum glucose and serum lipid lowering agents and are useful for the prophylaxis and treatment of diabetes, particularly type 2, and its complications, Syndrome X, the various forms of insulin resistance, and hyperlipidemias, and present reduced side effects, and, particularly, reduced or no liver toxicity.

REFERENCE COUNT: 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L20 ANSWER 3 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:511336 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 139:85372

TITLE: Preparation of pyrazolopyrimidines and related

compounds as hPPARlpha and hPPAR γ ligands

INVENTOR(S): Das, Saibal Kumar; Bhuniya, Debnath; Madhavan, Gurram

Ranga; Iqbal, Javed; Chakrabarti, Ranjan

PATENT ASSIGNEE(S): Reddy's Laboratories Ltd., India

SOURCE: PCT Int. Appl., 139 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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WO 2003053974
                                20030703
                                           WO 2002-IB5442
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             PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ,
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                                                                A 20011221 <--
PRIORITY APPLN. INFO.:
                                            WO 2002-IB5442
                                                                W 20021217 <--
OTHER SOURCE(S):
                       CASREACT 139:85372; MARPAT 139:85372
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GI

/ Structure 120 in file .gra /

Title compds. I [R1 = H, halo, OH, etc.; R2 = H, OH, halo, etc.; R3 = H, AΒ (un) substituted alkyl, cycloalkyl, etc.; Z = O, NR4; R4 = H, (un) substituted alkyl, aryl, etc.; Y = O, S, NR6, etc.; R6 = H, (un)substituted alkyl, aryl, etc.; Ar = (un)substituted aromatic, heteroarom., heterocyclic; G = 0, S; X = O, NHR5, (CH2)pO, etc.; R5 = H, (un)substituted alkyl, aryl, etc.; n = 1-4; p = 0-4; A = (un)substituted pyrazolopyrimidine, imidazolopyrimidine] and their pharmaceutically acceptable salts and formulations were prepared For example, O-alkylation of 5-ethyl-1,4-dihydro-1-methyl-3-propyl-7H-pyrazolo[4,3d]pyrimidin-7-one by chloroacetyl II, e.g., prepared from 4-aminothiophenol in 3-steps, followed by ester hydrolysis, afforded claimed pyrazolopyrimidine III in 5% yield. In hPPARlpha and hPPAR γ Luciferase ligand binding assays, 2examples of compds. I, e.g., pyrazolopyrimidine III, exhibited activity at 50 and 1 μM , resp. The test compds. also inhibited HMG CoA reductase (no data provided). Compds. I are claimed useful as antidiabetic, hypolipidemic, antiobesity and hypocholesterolemic agents.

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L20 ANSWER 4 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:319859 CAPLUS Full-text DOCUMENT NUMBER: 138:337836

TITLE: Long-chain, unsaturated, aromatic dicarboxylic acid

derivatives, their preparation, and therapeutic use for treatment of conditions mediated by peroxisome

proliferator-activated receptors (PPAR).

INVENTOR(S): Sauerberg, Per; Bury, Paul Stanley; Jeppesen, Lone;

Mogensen, John Patrick

PATENT ASSIGNEE(S): Novo Nordisk A/S, Den. SOURCE: PCT Int. Appl., 104 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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KIND
                                DATE
                                           APPLICATION NO.
                                                                   DATE
     PATENT NO.
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                                           _____
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                        A1 20030424 WO 2002-DK692
     WO 2003033453
                                                                  20021015 <--
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20040721 EP 2002-772084
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     EP 1438283
                         A1
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                              20041026 BR 2002-13253
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                                                                    20021015 <--
                        A2
                             20041228
     HU 2004001837
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    CN 1571766 A 20050126 CN 2002-820547
JP 2005505616 T 20050224 JP 2003-536195
US 20030109579 A1 20030612 US 2002-272613
                                                                   20021015 <--
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                                                                   20021016 <--
                        B2 20070522
     US 7220877
                                          IN 2004-CN771 20040415 <--
DK 2001-1524 A 20011017 <--
US 2001-330346P P 20011018 <--
WO 2002-DK692 W 20021015 <--
     IN 2004CN00771
                        A 20060113
PRIORITY APPLN. INFO.:
OTHER SOURCE(S): MARPAT 138:337836
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/ Structure 121 in file .gra /

GΙ

AΒ A novel class of dicarboxylic acid derivs., I, is disclosed [wherein: A = (un) substituted C1-3 alkylene, or A'O or A'S where A' is (un) substituted C1-3 alkylene; B = (un)substituted C1-3 alkylene, or OB' or SB' where B' is (un) substituted C1-3 alkylene; D, E = H, C1-6 alkyl, C3-6 cycloalkyl; L, M = O or S; T, U = C3-9 divalent, (un) substituted, unsatd. carbon chain; X, Y = (un) substituted arylene or heteroarylene; Z = (un) substituted arylene, heteroarylene, or divalent polycyclic ring system]. Also disclosed is the use of I in pharmaceutical compns., pharmaceutical compns. comprising I, and methods of treatment employing I and the compns. The present compds. may be useful (no data) in the treatment and/or prevention of conditions mediated by peroxisome proliferator-activated receptors (PPAR). For example, 1,4diiodobenzene was coupled with excess 2-penten-4-yn-1-ol in (iso-Pr)2NH in the presence of CuI and Pd(PPh3)4 at 60° , to give 55% (E,E)-5-[4-(5-hydroxypent-3en-1-ynyl)phenyl]pent-2-en-4-yn-1-ol. Mitsunobu reaction of this diol with (S)-2-ethoxy-3-(4-hydroxyphenyl)propionic acid Et ester using azodicarboxylic acid dipiperidide and PBu3 in THF gave 27% invention compound II. A total of 29 synthetic examples illustrate a variety of I, mostly sym. diacids and diesters, and mostly stereoisomeric, with all stereoisomers having (E) and (S) stereochem. at double bonds and chiral centers. Claims list a wide variety of sym. and asym. I, all named without stereochem. Claimed applications include treatment of type I and II diabetes, dyslipidemia, syndrome X and its conditions, cardiovascular diseases including atherosclerosis, and hypercholesterolemia.

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L20 ANSWER 5 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:242286 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 138:271396

TITLE: Preparation of 3-aryl-2-alkoxypropanoates from

3-aryl-2-oxopropanoates via ketalization and reduction

INVENTOR(S): Siripragada, Mahender Rao; Vanadanapu, Loka Appala

Purushotham; Mamillapalli, Ramabhadra Sarma; Gaddam,

ADDITONTION NO

ם דיי עם

Om Reddy

PATENT ASSIGNEE(S): Reddy's Laboratories Ltd., India

SOURCE: PCT Int. Appl., 12 pp.

CODEN: PIXXD2

KIND DATE

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	.ENT .	NO.			KIN	_	DATE		•	APPL 	ICAT	TON .	NO.		D	ATE 		
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			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	NO,	NZ,	OM,	PH,	
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											WO 2	002-	IB38	74	1	₩ 2	0020	919	<

OTHER SOURCE(S): CASREACT 138:271396; MARPAT 138:271396

4-HOC6H4CH2CH(OR1)CO2R2 (R1 = H, alkyl; R2 = alkyl), were prepared by converting 4-HOC6H4CH2COCO2H to 4-HOC6H4CH2C(OR1)2CO2R2 (variables as above) in the presence of acid followed by reduction in HOAc at 40-80 psi for 6-12 h. Thus, 4-hydroxyphenylpyruvic acid was stirred in EtOH at 10-15° for 3 h, at room temperature for 6 h, and at 50-60° for 4 h to give 4-HOC6H4CH2C(OEt)2CO2Et. The latter was hydrogenated in EtOH/HOAc over Rh/Al2O3 at 60 psi for 12 h to give 4-HOC6H4CH2CH(OEt)CO2Et.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L20 ANSWER 6 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:964313 CAPLUS Full-text DOCUMENT NUMBER: 138:55745

TITLE: Preparation of substituted 3-phenyl-2-alkoxypropanoic

acids and analogs as modulators of peroxisome proliferator activated receptors for treatment of

diabetes and related conditions

INVENTOR(S): Brooks, Dawn Alisa; Warshawsky, Alan M.;

Montrose-Rafezadeh, Chahrzad; Reifel-Miller, Anne; Prieto, Lourdes; Rojo, Isabel; Martin, Jose Alfredo;

Gonzales Garcia, Maria Rosario; Torrado, Alicia; Ferritto Crespo, Rafael; Lamas-Peteira, Carlos; Martin-Ortega Finger, Maria; Ardecky, Robert J.

PATENT ASSIGNEE(S): Eli Lilly and Company, USA; Ligand Pharmaceuticals

Incorporated

SOURCE: PCT Int. Appl., 458 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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OTHER S	OURCE	(S):			MARI	PAT	138:	5574.	5									

OTHER SOURCE(S): MARPAT 138:55745

GΙ

/ Structure 122 in file .gra /

Title compds. I [wherein Ar = (un)substituted aryl; Q = covalent bond, CH2, CH2CH2, CH2CH2CH2, or CH2CH2CH2CH2; W = (un)substituted (hetero)alkylene from 2-10 atoms in length in which 1 or more methylene groups have been replaced with CH=CH, C.tplbond.C, O, CO, NR7, NR7CO, C(=NOH), S, SO, SO2, or CHNR7R8; ring A is optionally substituted with up to 4 substituents in addition to R1; R1 = (CH2)nCH(OR2)(CH2)mE, CH=C(OR2)(CH2)mE, (CH2)nCHY(CH2)mE, or

CH=CY(CH2)mE; E = CO2R3, alkylnitrile, carboxamide, or (un)substituted sulfonamide, acylsulfonamide, or tetrazole; R2 = H, haloalkyl, COR4, CO2R4, CONR5R6, CSR4, CSOR4, CSNR5R6, or (un)substituted aliphatic group, aralkyl, or aryl; Y = O, CH2, CH2CH2, or CH=CH bonded ortho to R1 on ring A; R3-R8 = independently H or (un)substituted aliphatic group or aryl; m and n = 1independently 0-2; or pharmaceutically acceptable salts, hydrates, stereoisomers, or solvates thereof] were prepared by solution phase and solid phase synthetic methods as peroxisome proliferator activated receptor (PPAR) modulators (no data). For example, (S)-2-methoxy-3-hydroxyphenylpropanoic acid Et ester was treated with Ph triflimide to give the 4trifluoromethanesulfonyloxyphenyl derivative (97%). Substitution with propargyl alc. in the presence of PdCi2(PPh3)2 and TEA in DMF afforded the 4-(3-hydroxyprop-1-ynyl) phenyl intermediate (32%), which was coupled with 4phenylphenol using the Mitsunobu procedure to give II. Binding and cotransfection studies showed that many of the compds. of the invention are selective PPAR γ agonists or PPAR α /PPAR γ co-agonists (no data). Thus, I are useful for the treatment of hyperglycemia, dyslipidemia, Type I or II diabetes, hypertriglyceridemia, syndrome X, insulin resistance, heart failure, diabetic dyslipidemia, hyperlipidemia, hypercholesteremia, hypertension, obesity, anorexia bulimia, polycystic ovarian syndrome, anorexia nervosa, cardiovascular disease or other diseases where insulin resistance is a component (no data).

ΤI Preparation of substituted 3-phenyl-2-alkoxypropanoic acids and analogs as modulators of peroxisome proliferator activated receptors for treatment of diabetes and related conditions

ΡI	WO 2002100813 A	A2 20021219		
	PATENT NO.		APPLICATION NO.	DATE
ΡI	WO 2002100813	A2 20021219	WO 2002-US16950	
		A3 20031127		
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			DZ, EC, EE, ES, FI, GB,	
			JP, KE, KG, KP, KR, KZ,	
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		, US, UZ, VN, YU, ZA,	•	
	· · ·		SL, SZ, TZ, UG, ZM, ZW,	
			BE, CH, CY, DE, DK, ES,	
	·		SE, TR, BF, BJ, CF, CG,	CI, CM, GA,
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			CA 2002-2449256	
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			BR 2002-10190	
			CN 2002-811530	
	HU 2004000280		HU 2004-280	20020530 <
	HU 2004000280	A3 20060130		
	JP 2005509590			
		A 20060127		
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	ZA 2003008863	A 20050214		
	US 20050020684			20031201 <
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	MX 2003PA11201		MX 2003-PA11201	
	US 20070276138	A1 20071129	US 2006-637223	20061211 <

20010607 <--20020530 <--

US 2001-297144P P WO 2002-US16950 W

PRAI US 2001-297144P

US 2003-479262 A1 20031201

AΒ Title compds. I [wherein Ar = (un)substituted aryl; Q = covalent bond, CH2, CH2CH2, CH2CH2CH2, or CH2CH2CH2CH2; W = (un)substituted (hetero)alkylene from 2-10 atoms in length in which 1 or more methylene groups have been replaced with CH=CH, C.tplbond.C, O, CO, NR7, NR7CO, C(=NOH), S, SO, SO2, or CHNR7R8; ring A is optionally substituted with up to 4 substituents in addition to R1; R1 = (CH2) nCH(OR2) (CH2) mE, CH=C(OR2) (CH2) mE, (CH2) nCHY(CH2) mE, orCH=CY(CH2)mE; E = CO2R3, alkylnitrile, carboxamide, or (un)substituted sulfonamide, acvlsulfonamide, or tetrazole; R2 = H, haloalkyl, COR4, CO2R4, CONR5R6, CSR4, CSOR4, CSNR5R6, or (un)substituted aliphatic group, aralkyl, or aryl; Y = O, CH2, CH2CH2, or CH=CH bonded ortho to R1 on ring A; R3-R8 = independently H or (un) substituted aliphatic group or aryl; m and n = 1independently 0-2; or pharmaceutically acceptable salts, hydrates, stereoisomers, or solvates thereof] were prepared by solution phase and solid phase synthetic methods as peroxisome proliferator activated receptor (PPAR) modulators (no data). For example, (S)-2-methoxy-3-hydroxyphenylpropanoic acid Et ester was treated with Ph triflimide to give the 4trifluoromethanesulfonyloxyphenyl derivative (97%). Substitution with propargyl alc. in the presence of PdCl2(PPh3)2 and TEA in DMF afforded the 4-(3-hydroxyprop-1-ynyl)phenyl intermediate (32%), which was coupled with 4phenylphenol using the Mitsunobu procedure to give II. Binding and cotransfection studies showed that many of the compds. of the invention are selective PPARy agonists or PPAR α /PPARy co-agonists (no data). Thus, I are useful for the treatment of hyperglycemia, dyslipidemia, Type I or II diabetes, hypertriglyceridemia, syndrome X, insulin resistance, heart failure, diabetic dyslipidemia, hyperlipidemia, hypercholesteremia, hypertension, obesity, anorexia bulimia, polycystic ovarian syndrome, anorexia nervosa, cardiovascular disease or other diseases where insulin resistance is a component (no data).

ΙT 348-27-6P, 2-Fluoro-4-hydroxybenzaldehyde 405-05-0P, 3-Fluoro-4-hydroxybenzaldehyde 627-18-9P, 3-Bromopropan-1-ol 1073-05-8P, [1,3,2]Dioxathiane 2,2-dioxide 2973-78-6P, 3-Bromo-4-hydroxybenzaldehyde 3351-60-8P, 4-(2-Bromoethoxy)biphenyl 16251-33-5P, 1-Bromo-3-(4-phenoxyphenyl)propane 19070-95-2P, 2-(Biphenyl-4-yloxy)ethanol 23418-85-1P, Toluene-4-sulfonic acid but-3-ynyl ester 29169-19-5P 54334-74-6P, (Biphenyl-4-yloxy)acetic 63457-51-2P, 1-(3-Bromopropoxy)-4-phenoxybenzene acid ethyl ester 69455-12-5P, 4-Benzyloxy-3-bromobenzaldehyde 87545-48-0P, 96363-80-3P, Methanesulfonic acid 4-(2-Bromoethoxy) phenoxybenzene 3-dimethylaminopropyl ester 102229-10-7P, 2-(tert-Butyldimethylsilanyloxy)ethanol 111915-33-4P, 4-(2,2,3,3-Tetrafluoropropoxy)phenol 113795-28-1P, 4-(3-Bromopropoxy) biphenyl 119437-35-3P, 1-Chloro-3-(4-phenoxyphenyl)propane 128316-64-3P, 3-(4-Benzyloxyphenyl)-2-hydroxypropanoic acid methyl ester 156335-14-7P, Methyl 3-(4-hydroxyphenyl)-2-methoxypropanoate 156335-15-8P, 2-Ethoxy-3-(4-hydroxyphenyl)propionic acid methyl 156659-87-9P, (2S,4S)-4-(tert-Butyldimethylsilanyloxy)pentan-2-ol 173025-78-0P, 3-(Biphenyl-4-yloxy)propan-1-ol183612-97-7P, (1R*,3S*)-3-(tert-Butyldimethylsilanyloxy)cyclopentanol183795-20-2P, trans-3-(tert-Butyldimethylsilanyloxy)cyclopentanol 211617-68-4P 222835-03-2P, 3-(4-Benzyloxyphenyl)-2-ethoxyacrylic acid ethyl ester 223126-28-1P, 3-(4-Benzyloxyphenyl)-2-ethoxypropionic acid ethyl ester 251978-39-9P, 3-(4-Hydroxyphenyl)-2-phenoxypropanoic acid methyl ester 267228-40-0P, (S)-3-(4-Benzyloxyphenyl)-2hydroxypropionic acid ethyl ester 257228-41-1P, (2S)-2-Hydroxy-3-(4-hydroxyphenyl)propionic acid ethyl ester 325827-53-9F, (S)-3-(4-Hydroxyphenyl)-2-isopropoxypropionic acid 361576-28-5P, 3-(4-Benzyloxyphenyl)-2-ethoxy-3ethvl ester hydroxypropionic acid ethyl ester 477979-19-4P,

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(2S)-2-Methoxy-3-(4-trifluoromethanesulfonyloxyphenyl)propionic acid ethyl
ester 477979-21-8P, (2S)-3-[4-(3-Hydroxyprop-1-yny1)pheny1]-2-
methoxypropionic acid ethyl ester
                                  477979-26-3P,
(2S)-3-[4-(3-Chloroprop-1-yny1)pheny1]-2-methoxypropionic acid ethyl ester
477979-44-5P, (2S)-3-[4-(5-Hydroxypent-1-ynyl)phenyl]-2-methoxypropionic
acid ethyl ester 477979-49-0P, 3-[4-(5-Bromopent-1-ynyl)phenyl]-2-
methoxypropionic acid ethyl ester
                                   477979-66-1P, 4-But-3-ynyloxybiphenyl
477979-67-2P, (2S)-3-[4-[4-(Biphenyl-4-yloxy)] but -1-ynyl] phenyl] -2-
methoxypropionic acid ethyl ester 477979-69-4P,
1-(But-3-ynyloxy)-4-phenoxybenzene 477979-71-8P,
[4-(But-3-ynyloxy)phenyl]phenylmethanone
                                          477979-72-9P,
(2S)-3-[4-[4-(4-Benzoylphenoxy)but-1-ynyl]phenyl]-2-methoxypropionic acid
             477979-80-9P, (2S)-3-[4-(6-Hydroxyhex-1-ynyl)phenyl]-2-
ethvl ester
methoxypropionic acid ethyl ester
                                   477979-88-7P,
(2S)-3-[4-[4-(4-Benzoylphenoxy)butyryl]phenyl]-2-methoxypropionic acid
             477979-96-7P, cis-2-(tert-
Butyldimethylsilanyloxy)cyclopentanol
                                       477979-97-8P
                                                      477979-99-0P
477980-00-0P 477980-11-3P, (2R,3S)-3-(4-Phenoxyphenoxy) butan-2-ol
              477980-13-5P 477980-20-4P,
477980-12-4P
(2S)-3-(3'-Hydroxymethylbiphenyl-4-yl)-2-methoxypropionic acid ethyl ester
477980-23-7P 477980-24-8P 477980-35-1P 477980-36-2P,
3-(Biphenyl-4-yloxy)cyclohexanol
                                  477980-37-3P,
(trans)-3-(Biphenyl-4-yloxy)cyclohexanol
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(cis)-3-(Biphenyl-4-yloxy)cyclohexanol
                                        477980-44-2P,
(2S)-3-[4-(tert-Butyldimethylsilanyloxy)phenyl]-2-methoxypropionic acid
477980-45-3P, (2S)-3-[4-(3-Hydroxypropoxy)phenyl]-2-methoxypropionic acid
477980-56-6P, (2S)-3-[4-(2-Hydroxyethoxy)phenyl]-2-methoxypropanoic acid
477980-59-9P, (2S)-3-(4-Ethynylphenyl)-2-methoxypropionic acid ethyl ester
477980-60-2P, (2S)-3-(4-Acetylphenyl)-2-methoxypropionic acid ethyl ester
477980-61-3P, (2S)-3-[4-(2-Bromoacetyl)phenyl]-2-methoxypropionic acid
ethyl ester 477980-66-8P, (2S)-3-[4-(4-Hydroxybutyl)phenyl]-2-
methoxypropionic acid ethyl ester 477980-69-1P,
3-(4-Benzyloxy-3-methoxyphenyl)-3-hydroxy-2-methoxypropionic acid methyl
ester 477980-70-4P, 3-(4-Hydroxy-3-methoxyphenyl)-2-
methoxypropionic acid methyl ester
                                    477980-71-5P,
3-(4-Hydroxy-3-methoxyphenyl)-2-methoxypropionic acid 477980-72-6P,
3-[4-(tert-Butyldimethylsilanyloxy)-3-methoxyphenyl]-2-methoxypropionic
acid 477980-76-0P, 3-(4-Hydroxy-3-methoxyphenyl)-2-
methoxypropionic acid ethyl ester 477980-77-1P,
3-[4-[3-(Biphenyl-4-yloxy)propoxy]-3-methoxyphenyl]-2-methoxypropionic
acid ethyl ester 477980-80-6P,
(2S)-3-(3-Chloro-4-hydroxyphenyl)-2-methoxypropionic acid ethyl ester
477980-82-8P, (2S)-3-[4-[3-(Biphenyl-4-yloxy)propoxy]-3-chlorophenyl]-2-
methoxypropionic acid ethyl ester 477980-84-0P,
                                             477980-85-1P,
2-(3-Fluoro-4-methoxyphenyl)-[1,3]dioxolane
4-[1,3]Dioxolan-2-yl-2-fluorophenol 477980-86-2P,
4-[3-(Biphenyl-4-yloxy)propoxy]-3-fluorobenzaldehyde
3-[4-[3-(Biphenyl-4-yloxy)propoxy]-3-fluorophenyl]-3-hydroxy-2-
methoxypropionic acid methyl ester 477980-88-4P,
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methvl ester
              477980-90-8P, 4-Benzyloxy-3-trifluoromethylbenzaldehyde
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methyl ester
trifluoromethylphenyl]-2-methoxyacrylic acid methyl ester 477980-93-1P,
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methoxyacrylic acid 477980-95-3P 477980-96-4P,
(2S)-3-(6-Hydroxy-4'-methoxybiphenyl-3-yl)-2-methoxypropionic acid ethyl
       477980-97-5P, (2S)-3-[6-[3-(Biphenyl-4-yloxy)propoxy]-4'-
methoxybiphenyl-3-yl]-2-methoxypropionic acid ethyl ester 477981-00-3P,
2-Methyl-4-(triisopropylsilanyloxy)benzaldehyde 477981-01-4P,
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3-[4-[3-(Bipheny1-4-yloxy)propoxy]-2-methylphenyl]-2-methoxyacrylic acid
methyl ester 477981-02-5P, 3-(4-Hydroxy-2-methylphenyl)-2-methoxyacrylic
          477981-04-7P, 3-[4-[3-(Biphenyl-4-yloxy)propoxy]-2-methylphenyl]-2-
methoxypropionic acid methyl ester 477981-07-0P,
3-(3-Hydroxyphenyl)-2-methoxypropionic acid methyl ester
                                                                                         477981-20-7P,
(2S)-3-[4-(3-Bromopropoxy)phenyl]-2-methoxypropionic acid ethyl ester
477981-27-4P, (S)-5-(4-Benzyloxybenzyl)-2,2-dimethyl-[1,3]dioxolan-4-one
477981-33-2P, (S)-5-(4-Hydroxybenzyl)-2,2-dimethyl-[1,3]dioxolan-4-one
477982-20-0P, 4-[3-(Biphenyl-4-yloxy)propoxy]-2-fluorobenzaldehyde
477982-21-1P, 3-[4-[3-(Biphenyl-4-yloxy)propoxy]-2-fluorophenyl]-3-hydroxy-
2-methoxypropionic acid methyl ester
                                                          477982-22-2P,
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                      477982-24-4P, 3-(4-Benzyloxyphenyl)-2-(4-
methvl ester
chlorophenoxy) propanoic acid methyl ester
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2-Phenoxy-3-[4-[3-(4-phenoxyphenoxy)propoxy]phenyl]propanoic acid methyl
            477982-27-7P, Methyl 3-hydroxy-2-methoxy-3-[4-
(phenylmethoxy)phenyl]propanoate
                                                    477982-28-8P,
3-(4-Hydroxyphenyl)-2-methoxypropanoic acid 477982-29-9P
                                                                                             477982-30-2P,
Ethyl (2S)-2-methoxy-3-[4-[3-(4-phenoxyphenoxy)propoxy]phenyl]propanoate
477982-37-9P, (2S)-2-Hydroxy-3-[4-[3-(4-
phenoxyphenoxy)propoxy]phenyl]propionic acid ethyl ester 477982-38-0P,
(2S)-2-Ethoxy-3-[4-[3-(4-phenoxyphenoxy)propoxy]phenyl]propionic acid
                    477982-41-5P, (2S)-3-[4-[3-[4-(4-
Hydroxybenzoy1)phenoxy[propoxy[pheny1]-2-methoxypropionic acid ethyl ester
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Butyldimethylsilanyloxy)ethoxy]benzoyl]phenoxy]propoxy]phenyl]-2-
                                                    477982-45-9P,
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(2S)-2-Allyloxy-3-(4-benzyloxyphenyl)propionic acid ethyl ester
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ethyl ester 477982-47-1P, (2S)-3-[4-[3-(4-Phenoxyphenoxy)propoxy]phenyl]-
2-propoxypropionic acid ethyl ester 477982-49-3P,
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(2S)-3-(3-Chloro-4-hydroxyphenyl)-2-ethoxypropionic acid ethyl ester
477982-54-0P, (2S)-3-[4-[3-(4-Benzoylphenoxy)propoxy]-3-chlorophenyl]-2-
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ester
methoxycarbonylethyl)phenoxy]propoxy]biphenyl-4-carboxylic acid methyl
           477982-59-5P, 3-[4-(3-Bromopropoxy)-2-methoxy-phenyl]-2-
methoxypropionic acid methyl ester 477982-61-9P,
(2S)-3-[4-[3-(4'-tert-Butylbiphenyl-4-yloxy)propoxy]-2-methoxyphenyl]-2-
methoxypropionic acid methyl ester 477982-64-2P,
4-(2,2,3,3-Tetrafluoropropoxy)1-benzyloxybenzene
                                                                             477982-65-3P,
(2S)-2-Methoxy-3-[4-[3-[4-(2,2,3,3-
tetrafluoropropoxy)phenoxy]propoxy]phenyl]propionic acid ethyl ester
477982-68-6P, (2S)-3-[4-[3-(4-Benzyloxyphenoxy)propoxy]pheny1]-2-
methoxypropionic acid ethyl ester
                                                      477982-69-7P,
(2S)-3-[4-[3-(4-Hydroxyphenoxy)propoxy]phenyl]-2-methoxypropionic acid
ethyl ester
                     477982-70-0P, (2S)-2-Methoxy-3-[4-[3-[4-(3-
methylbutoxy)phenoxy]propoxy]phenyl]propionic acid ethyl ester
477982-78-8P, (2S)-3-[4-[3-(4-Iodophenoxy)propoxy]pheny1]-2-
methoxypropionic acid ethyl ester
                                                    477982-79-9P,
(2S)-3-[4-[3-[4-(1H-Indol-5-y1)phenoxy]propoxy]pheny1]-2-methoxypropionic
acid ethyl ester 477982-84-6P, (2S)-3-[4-[3-(4'-Cyanobiphenyl-4-
yloxy)propoxy]phenyl]-2-methoxypropionic acid ethyl ester
                                                                                           477982-86-8P,
(2S)-2-Methoxy-3-[4-[3-[4'-(1H-tetrazol-5-yl)biphenyl-4-
yloxy]propoxy]phenyl]propionic acid ethyl ester 477982-91-5P,
(2S)-2-Methoxy-3-[4-[3-[4-(piperazin-1-yl)phenoxy]propoxy]phenyl]propionic
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acid ethyl ester 477982-93-7P, (2S)-2-Methoxy-3-[4-[3-[4-(morpholin-4yl)phenoxy]propoxy]phenyl]propionic acid ethyl ester 477982-95-9P, 3-[4-[3-(Biphenyl-4-yloxy)propoxy]-2-chlorophenyl]-2-hydroxypropionic acid 477982-97-1P, (2S)-3-[4-(2-Bromoethoxy)phenyl]-2-methoxypropionic acid477983-02-1P, 3-(3-Benzyloxyphenyl)-3-hydroxy-2ethyl ester methoxypropionic acid methyl ester 477983-03-2P, 3-(3-Benzyloxyphenyl)-2-methoxyacrylic acid methyl ester 477983-04-3P 477983-05-4P, 3-(3-Benzyloxyphenyl)-2-methoxypropionic acid methyl ester477983-44-1P, 3-[3-(3-Bromopropoxy)phenyl]-2-methoxypropionic acid methyl 477983-82-7P, 3-[3-(2-Bromoethoxy)phenyl]-2-methoxypropionic acid 477984-10-4P, (2S)-3-(4-Benzyloxyphenyl)-2-propoxypropionicmethyl ester 477984-12-6P, 2-Ethoxy-3-(3-hydroxyphenyl) propionic acid ethyl ester acid methyl ester

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation); RACT (Reactant or reagent)

(intermediate; preparation of substituted (phenyl)(alkoxy)propanoic acids and analogs as PPAR modulators for treatment of diabetes and related conditions)

IT 186895-45-4P, 3-(4-Benzyloxyphenyl)propionic acid ethyl ester 477980-81-7P, (2S)-3-(3,5-Dichloro-4-hydroxyphenyl)-2-methoxypropionic acid ethyl ester

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of substituted (phenyl)(alkoxy)propanoic acids and analogs as PPAR modulators for treatment of diabetes and related conditions)

L20 ANSWER 7 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:793403 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 137:310931

TITLE: Preparation of phenylalkanoic acid derivatives as preventive or remedial agents for digestive tract

diseases

INVENTOR(S): Horizoe, Tatsuo; Shinoda, Masanobu; Emori, Eita;

Matsuura, Fumiyoshi; Kaneko, Toshihiko; Ohi, Norihito; Kasai, Shunji; Yoshitomi, Hideki; Yamazaki, Kazuto; Miyashita, Sadakazu; Hihara, Taro; Seiki, Takashi;

Clark, Richard; Harada, Hitoshi

PATENT ASSIGNEE(S): Eisai Co., Ltd., Japan SOURCE: PCT Int. Appl., 344 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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A	AU 2002242989 A1 20021021								1021		AU 2	002-	24298	39		20	0020	327 <	<
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/ Structure 123 in file .gra /

AΒ Disclosed is a preventive/remedy for digestive tract or inflammatory diseases, which contains as the active ingredient a novel carboxylic acid derivative represented by the following formula [I; R1 = H, OH, each (un)substituted C1-6alkyl, C1-6 alkoxy, C1-6 alkylthio, C1-6 hydroxyalkyl, C1-6 hydroxyalkoxy, C1-6 hydroxyalkylthio, C1-6 aminoalkyl, C1-6 aminoalkoxy, C1-6 aminoalkylthio, C2-12 alkoxyalkyl, C3-7 cycloalkyl, C3-7 cycloalkyloxy, C3-7 cycloalkylthio, C2-6 alkenyl, C2-6 alkenyloxy, or C2-6 alkenylthio, etc.; L = a single or double bond, each (un)substituted C1-6 alkylene, C2-6 alkenylene, or C2-6 alkynylene; M = a single bond, each (un)substituted C1-6 alkylene, C2-6 alkenylene, or C2-6 alkynylene; T = a single bond, each (un)substituted C1-3alkylene, C2-3 alkenylene, or C2-3 alkynylene; W = 2,4-dioxothiazolidin-5-yl,2,4-dioxothiazolidin-5-ylidene, carboxy, (un) substituted CONH2; X=0, (un) substituted C2-6 alkenylene, hydroxymethylene, CO, CS, N-(un) substituted CQNH, NHCQ, SO2NH, NHSO2, or NHCQNH (Q = 0, S); Y = (un) substituted C5-12 aromatic hydrocarbyl or C3-7 aliphatic hydrocarbyl optionally containing ≥1 heteroatoms; ring Z = C5-6 aromatic hydrocarbyl; Y = (un) substituted aromatic hydrocarbon group optionally containing ≥1 heteroatoms; some provisos given], a salt of the derivative, or a hydrate of either. The above digestive tract diseases include (1) inflammatory digestive tract diseases such as ulcerous colitis, Crohn's disease, pancreatitis, and gastritis, (2) digestive tract proliferative diseases such as digestive tract benign rumors, digestive tract polyp, hereditary (genetic) polyposis syndromes, colon cancer, rectum cancer, and stomach cancer, and (3) digestive tract ulcerous diseases such as duodenal ulcer, stomach ulcer, esophagus ulcer, regurgitant esophagitis, stress ulcer or erosion, erosion caused by drugs, and Zollinger-Ellison syndromes. The above inflammatory diseases include arthritic rheumatism, multiple sclerosis, immunodeficiency, cachexia, osteoarthritis, osteoporosis, asthma, and allergy. The compds. I are triple agonists for PPAR (peroxisome proliferator-activated receptor) α , β , and γ subtype. Thus,

2-isopropoxy-3-[4-methoxy-3-[[[4-

(trifluoromethyl)benzyl]amino]carbonyl]phenyl]propanoic acid in vitro showed the transcription activity for PPARa, β , and γ with EC50 of 0.08, 2.513, and 0.382 μM , resp., in CV-1 cell. (2S)-3-[3-[[(2,4-dichlorobenzoyl)amino]methyl]-4-methoxyphenyl]-2- isopropoxypropanoic acid at 1 mg/kg/day p.o. for 3 days showed a disease activity index based on diarrhea, bloody excrement, and weight loss (DAI) of 2.0±0.3 in mice suffering from colitis induced by dextran sulfate sodium salt vs. 2.8±0.2 for the control group and 2.1±0.3 for the mice treated with rosiglitazone at 30 mg/kg/day. Many compds. prepared do not possess the thiazolidine skeleton and thereby may completely avoid toxicity such as liver disorder which was noted in the past as a problem for compds. having PPARy agonist activity.

L20 ANSWER 8 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:286703 CAPLUS Full-text

DOCUMENT NUMBER: 136:309930

TITLE: Preparation of benzimidazole derivatives for treatment

and prevention of diabetes

INVENTOR(S): Fujita, Takashi; Wada, Kunio; Koguchi, Minoru; Honma,

Eiji; Fujiwara, Toshihiko

PATENT ASSIGNEE(S): Sankyo Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 135 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GT

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2002114781	A	20020416	JP 2000-307157	20001006 <
PRIORITY APPLN. INFO.:			JP 2000-307157	20001006 <
OTHER SOURCE(S):	MARPAT	136:309930		

/ Structure 124 in file .gra /

AB The title compds. I [R1 - R6 = H, alkyl, etc.; n, q = 1 - 5; Q, Y = 0, S; X = CH2, etc.; Z = CH, N; A = (CH2)mCH(CO2H)BR7, etc.; B = 0, etc.; R7 = H, alkyl, etc.; m = 0 - 8] are prepared Compds. of this invention at 0.01% in feed (given for 3 days) gave 34.9% to 66.7% decrease of blood sugar in diabetic KK mice.

TI Preparation of benzimidazole derivatives for treatment and prevention of diabetes

PI JP 2002114781 A 20020416

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	JP 2002114781	A	20020416	JP 2000-307157	20001006 <
PRAI	JP 2000-307157		20001006	<	

IT Arteriosclerosis

Arthritis

Autoimmune disease

Cachexia

Cardiovascular system, disease

Glaucoma (disease)

Gout

Hypertension

Osteoporosis

Rheumatoid arthritis

(preparation and effect of benzimidazole derivs.)

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation); RACT (Reactant or reagent)

(preparation of benzimidazole derivs. for treatment and prevention of diabetes)

L20 ANSWER 9 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:157746 CAPLUS Full-text

DOCUMENT NUMBER: 136:200176
TITLE: Preparation of

3-[(oxazolylalkoxy)phenyl]-2-phenoxypropionic acid derivatives as PPAR agonists for treatment of diabetes

mellitus and related conditions

INVENTOR(S): Ardecky, Robert J.; Brooks, Dawn Alisa; Godfrey,

Alexander Glenn; Jones, Sarah Beth; Mantlo, Nathan Bryan; McCarthy, James Ray; Michellys, Pierre-Yves; Rito, Christopher John; Tyhonas, John S.; Winneroski,

Leonard Larry; Xu, Yanping

PATENT ASSIGNEE(S): Eli Lilly and Company, USA; Ligand Pharmaceuticals

SOURCE: PCT Int. Appl., 217 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GΙ

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		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KΡ,	KR,	KΖ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	NO,	NZ,	PH,	PL,
		PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,
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							GB,										
		ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML ,	MR,	ΝE,	SN,	TD,	ΤG	
CA	2418	134			A1		2002	0228		CA 2	001-	2418	134		2	0010	823 <
AU	2001	0846	60		Α		2002	0304		AU 2	001-	8466	0		2	0010	823 <
EP	1313	717			A1		2003	0528		EP 2	001-	9637	34		2	0010	823 <
EP	1313	717			В1		2007	1017									
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	ΑL,	TR						
JP	2004	5067	22		T		2004	0304		JP 2	002-	5214	33		2	0010	823 <
AT	3759	85			T		2007	1115		AT 2	001-	9637	34		2	0010	823 <
ES	2295	200			Т3		2008	0416		ES 2	001-	9637	34		2	0010	823 <
МX	2003	PA01	610		A		2003	0910		MX 2	003-	PA16	10		2	0030	221 <
US	2004	0138	277		A1		2004	0715		US 2	003-	3431	87		2	0030	729 <
US	7176	224			В2		2007	0213									
PRIORITY	Y APP	LN.	INFO	.:						US 2	000-	2274	56P		P 2	0000	823 <
										WO 2	001-	US22	617	Ī	W 2	0010	823 <
OTHER SO	OURCE	(S):			MARI	PAT	136:	2001	76								

^{*} STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. I [wherein n = 2-4; R1 = H, (halo)alkyl, or Ph; R2 and R3 = independently H, alkyl, cycloalkyl(alkyl), alkoxy, or aryl(alkyl); or R2 forms (tetrahydro)naphthyl together with the Ph to which they are bound; R4 = alkyl; R5 = independently H or (un)substituted (hetero)aryl, with provisos; R6 = H or (amino)alkyl; R7 and R8 = independently H, (cyclo)alkyl, (halo)alkoxy, or halo(alkyl); or R8 form benzodioxolyl together with the Ph to which they are bound; and pharmaceutically acceptable salts, solvates, and hydrates thereof] were prepared as agonists of peroxisome proliferator activated receptors

(PPARs). For example, 2-[2-(3-bromopheny1)-5-methyloxazol-4-yl] ethanol was coupled with p-fluorophenyl boronic acid in the presence of PPh3, Pd(OAc)2, and Na2CO3 to give the biphenyl derivative (36%). Esterification with tosyl anhydride in the presence of pyridine and DMAP, followed by reaction with 3-(4-hydroxyphenyl)-2-methyl-2-phenoxypropionic acid Et ester in the presence of polystyrene bound 1,5,7-triazabicyclo[4.4.0]dec-5-ene and hydrolysis with NaOH, afforded II (24%). The latter bound to PPAR α and PPAR γ with IC50 values of 147 nM and 41 nM, resp., and activated the nuclear transcription factors huPPAR α and huPPAR γ with cotransfection efficacies of 38% and 93%, resp. In addition, HDLc serum levels increased by 40.4% in male transgenic mice dosed with 30 mg/kg of II, and glucose levels were normalized to 91% in male diabetic (db/db) mice dosed with 30 mg/kg of II. Thus, I are useful in the treatment and prevention of diabetes mellitus and related conditions.

L20 ANSWER 10 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2001:115318 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 134:177470

TITLE: Process for the preparation of substituted

3-phenyl-propanoic acid esters and substituted

3-phenyl-propanoic acids

INVENTOR(S): Ebdrup, Soren; Deussen, Heinz-Josef W.; Zundel,

Magali; Bury, Paul Stanley

PATENT ASSIGNEE(S): Novo Nordisk A/S, Den. SOURCE: PCT Int. Appl., 118 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

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PATENT NO.
                 KIND
                         DATE APPLICATION NO.
                  A1 20010215 WO 2000-DK440
WO 2001011073
                                                           20000807 <--
   W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
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       HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
       LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
       SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU,
       ZA, ZW
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       CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
EP 1206565
                   A1 20020522 EP 2000-952953
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JP 2003506065
                   Т
                         20030218
                                    JP 2001-515321
                                                           20000807 <--
WO 2002012472
                         20020214
                                     WO 2001-DK508
                                                           20010719 <--
                   A1
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       GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
       LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
       RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US,
       UZ, VN, YU, ZA, ZW
   RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
       DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
       BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
AU 2001081739 A 20020218 AU 2001-81739 20010719 <--
EP 1309674
                   A1
                         20030514 EP 2001-960183 20010719 <--
                  В1
                         20070613
EP 1309674
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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                                            AT 2001-960183
     AT 364691
                         T
                               20070715
                                                                    20010719 <--
     ES 2288976
                         Т3
                                20080201
                                            ES 2001-960183
                                                                    20010719 <--
                        A1 20030109
A1 20031023
B2 20060815
     US 20030008361
                                            US 2002-132428
                                                                     20020424 <--
     US 20030199048
                                            US 2003-343879
                                                                     20030205 <--
                                            US 1999-1101 A 19990805 <--
US 1999-148643P P 19990812 <--
US 2000-633613 B1 20000807 <--
WO 2000-DK439 A 20000807 <--
WO 2000-DK440 W 20000807 <--
DK 2001-88 A 20010117
US 2001-26336
     US 7091023
PRIORITY APPLN. INFO.:
                                             US 2001-263364P P 20010123 <--
WO 2001-DK508 W 20010719 <--
OTHER SOURCE(S):
                         CASREACT 134:177470; MARPAT 134:177470
     A process is provided for the preparation of optically enriched substituted
     esters of 3-phenyl-propanoic acids and substituted 3-phenyl-propanoic acids by
     the hydrolysis or transesterification of one of the two enantiomeric forms of
     a racemic or enantiomerically enriched 3-phenyl-propanoic acid ester by an
     enzyme. This enzymic resolution may be catalyzed by a large number of com.
     available lipases, proteases, peptidases, esterases or other hydrolytic
     enzymes. Thus, Et (2RS)-2-ethoxy-3-(4-hydroxyphenyl)propanoic acid ester was
     converted to (2S)-2-ethoxy-3-(4-hydroxyphenyl)propanoic acid and Et (2R)-2-
     ethoxy-3-(4-hydroxyphenyl)propanoic acid ester by immobilized Mucor mihei
     lipase in 9 h with an enantiomeric selectivity of 81% for the ester.
=> d 120 ibib abs ti hit 11-15
L20 ANSWER 11 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2000:742095 CAPLUS Full-text
DOCUMENT NUMBER:
                         133:296438
TITLE:
                         Preparation of substituted fused imidazole derivatives
                         as hypoglycemics
                         Fujita, Takashi; Wada, Kunio; Oguchi, Minoru; Honma,
INVENTOR(S):
                         Hidehito; Fujiwara, Toshihiko
PATENT ASSIGNEE(S):
                         Sankyo Company, Ltd., Japan
                         PCT Int. Appl., 274 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
     PATENT NO.
                        KIND DATE
                                           APPLICATION NO.
                                                                    DATE
                                _____
                                             _____
                         ____
                                                                     _____
     WO 2000061582
                         A1 20001019 WO 2000-JP2217 20000406 <--
         W: AU, BR, CA, CN, CZ, HU, ID, IL, IN, KR, MX, NO, NZ, PL, RU, TR,
         RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE
     JP 2000351777
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                                             JP 2000-105985
                                                                     20000407 <--
                         Α
                                             JP 1999-101369 A 19990408 <--
PRIORITY APPLN. INFO.:
```

OTHER SOURCE(S): MARPAT 133:296438

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AΒ Compds. represented by general formula (I) and salts and esters thereof [wherein R1 is hydrogen, C1-6 alkyl, (un)substituted C6-10 aryl or C7-16 aralkyl, HO, (un) substituted acyloxy, C1-6 alkoxy, (un) substituted NH2, etc.; R2 is hydrogen, C1-6 alkyl, or (un)substituted C7-16 aralkyl; R4, R4, or R5 is each hydrogen, C1-6 alkyl, or C1-6 alkoxy; R6 is hydrogen, C1-6 alkyl, (un) substituted C6-10 aryl or C7-16 aralkyl; Q and Y are each oxygen or sulfur; X is CH2, CO, CH(OR9), or C(:NOR10); wherein R9 or R10 is hydrogen, (un) substituted C1-6 alkyl, C7-16 aralkyl, or acyl; Z is CH or nitrogen; n and q are each 1 to 5; and A is a group represented by general formula Q1, Q2, Q3, or (CH2)m CH(CO2H)-BR7; wherein m is 0 to 8; X1 is oxygen or sulfur; B is oxygen, sulfur, or (un) substituted NH; and R7 is hydrogen, C1-6 alkyl, (un)substituted C6-10 aryl or C7-16 aralkyl, or haloalkyl] are prepared These compds. are useful as insulin resistance improvers, hypoglycemics, antiinflammatory agents, immunomodulators, aldose reductase inhibitors, 5lipoxygenase inhibitors, lipid peroxide-formation inhibitors, peroxisome proliferator-activated receptor (PPAR) activators, anti-osteoporosis agents, leukotriene antagonists, promoters of fat cell formation, cancer cellproliferation inhibitors, or calcium antagonists. They are useful for the prevention or treatment of diabetes, hyperlipidemia, obesity, glucose tolerance insufficiency, hypertension, fatty liver, diabetes complication, arteriosclerosis, gestational diabetes, polycystic ovarian syndrome, cardiovascular diseases, cell damages caused by atherosclerosis or ischemic heart diseases, gout, osteoarthritis, rheumatic arthritis, allergic diseases, asthma, gastrointestinal ulcer, cachexia, autoimmune diseases, cancer, osteoporosis, or cataract. Thus, N-[2-amino-5-(6-methoxymethoxy-2,5,7,8tetramethylchroman-2- ylmethoxy)phenyl]-N-methylcarbamic acid tert-Bu ester was condensed with 4-(2,4-dioxothiazolin-5-ylmethyl)phenoxyacetic acid using di-Et cyanophosphate and Et3N in THF at room temperature for 30 min, followed by treatment of the product with 4 N HCl/dioxane at room temperature for 5 h gave 5-[4-[6-(6-hydroxy-2,5,7,8-tetramethylchroman-2-ylmethoxy)-1-methyl-1Hbenzimidazol-2-ylmethoxy]benzyl]thiazolidine-2,4-dione hydrochloride (II.HCl). When a feed containing 0.01% II.HCl was fed to mice for 3 days, the blood sugar level was lowered by 66.7% compared to control animal.

L20 ANSWER 12 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1999:383506 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 131:179289

TITLE: Non-thiazolidinedione antihyperglycemic agents. Part

3: The effects of stereochemistry on the potency of

 α -methoxy- β -phenylpropanoic acids

AUTHOR(S): Haigh, David; Allen, Graham; Birrell, Helen C.;

Buckle, Derek R.; Cantello, Barrie C. C.; Eggleston, Drake S.; Haltiwanger, R. Curtis; Holder, Julie C.; Lister, Carolyn A.; Pinto, Ivan L.; Rami, Harshad K.; Sime, John T.; Smith, Stephen A.; Sweeney, John D.

CORPORATE SOURCE: SmithKline Beecham Pharmaceuticals, Essex, CM19 5AW,

UK

SOURCE: Bioorganic & Medicinal Chemistry (1999),

7(5), 821-830

CODEN: BMECEP; ISSN: 0968-0896

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal

LANGUAGE: English

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/ Structure 126 in file .gra /

AB Rhizopus delemar lipase catalyzed ester hydrolysis of the α -methoxy- β -phenylpropanoate (I) affords the (R)-(+) and (S)-(-) isomers in > 84% enantiomeric excess. Absolute stereochem, was determined by a single crystal X-ray anal, of a related synthetic analog. The activity of these two enantiomers on glucose transport in vitro and as anti-diabetic agents in vivo is reported and their unexpected equivalence attributed to an enzyme-mediated stereospecific isomerization of the (R)-(+) isomer. Binding studies using recombinant human PPARy (peroxisomal proliferator activated receptor γ), now established as a mol. target for this compound class, indicate a 20-fold higher binding affinity for the (S) antipode relative to the (R) antipode.

L20 ANSWER 13 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1999:244628 CAPLUS Full-text

DOCUMENT NUMBER: 130:296612

TITLE: Preparation of amidocarboxylic acid derivatives as

inhibitors of aldose reductase, 5-lipoxygenase, and

lipid peroxide formation and as peroxisome

proliferator-activated receptor (PPAR) activators INVENTOR(S): Yanagisawa, Hiroaki; Sakurai, Mitsuya; Takamura,

Makoto; Fujiwara, Toshihiko

PATENT ASSIGNEE(S): Sankyo Company, Ltd., Japan

SOURCE: PCT Int. Appl., 720 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	TENT	NO.			KINI)	DATE									DATE		
WO	9918 W:									WO 1	1998- , KR,	JP43	96					
	RW:	AT, PT,		CH,	CY,	DE,	DK,	ES,	FI,	FR,	, GB,	GR,	IE,	IT,	LU	, MC,	NL,	
CA	2305	808			A1		1999	0415		CA 1	1998-	2305	808			19980	930	<
AU	9892	798			A		1999	0427		AU 1	1998-	9279	8			19980	930	<
AU	7381	34			В2		2001	0906										
EP	1026	149			A1		2000	0809		EP 3	1998-	9455	27			19980	930	<
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	2000				A3		2001											
RU	2176	999			C2		2001	1220			2000-					19980		
	6528							0304			2000-	-				20000		
ИО	2000	0016	89		А		2000	0531		NO 2	2000-	1689				20000		
	2000				А		2001				2000-					20000		
US	2004	0006	141		A1		2004	0108			2002-					20020		
PRIORIT	Y APP	LN.	INFO	.:							1997–					19971		
											1998-					19980		
										US 2	2000-	5407	65		А3	20000	330	<

GΙ

/ Structure 127 in file .gra /

AΒ Claimed and prepared are amidocarboxylic acid derivs. (phenylalkanoic acids containing arylcarboxamide derivs.) represented by general formula (I), pharmacol. acceptable salts thereof, or pharmacol. acceptable esters thereof, [wherein R1 = H, linear or branched C1-6 alkyl, C7-12 aralkyl; R2 = linear or branched C1-6 alkylene; R3 = H, linear or branched alkyl C1-6 alkyl, C1-4 alkoxy, or C1-4 alkylthio, halo, NO2, di(linear or branched C1-4 alkyl)amino, (un) substituted C6-10 aryl, C7-12 aralkyl optionally having 1-5 substituents on the aryl, OH, linear or branched C1-5 aliphatic acyl; R4 = H, linear or branched C1-6 alkyl; Z = linear or branched C1-6 alkylene; W = HO, linear or branched C1-6 alkyl, C1-4 alkoxy, or C1-4 alkylthio, (un)substituted C6-10 aryl, C6-10 aryloxy, C6-10 arylthio, C7-12 aralkyloxy, C7-12 aralkylthio, or C6-10 aryloxy-linear or branched C1-4 alkyl each optionally having 1-5 substituents on the aryl, 5- to 10-membered mono- or bicyclic heteroaryloxy containing 1-4 heteroatoms selected from O, N, and S, etc.; X = C6-10 aryl optionally having 1-3 substituents, 5- to 10-membered mono- or bicyclic heteroaryl containing 1-4 heteroatoms selected from O, N, and S; Y = single bond, O, S, (un) substituted NH]. Also claimed are blood sugar- and blood lipid-lowering agents, insulin resistance improver, antiinflammatory agents, immunomodulators, aldose reductase inhibitors, 5-lipoxygenase inhibitors, lipid peroxide formation inhibitors, PPAR activators, and anti-osteoporosis agents and therapeutic or prophylactic agents for diabetes, hyperlipemia, obesity, impaired glucose tolerance, insulin resistant non-impaired glucose tolerance, fatty liver, diabetes complications, gestational diabetes mellitus, polycystic ovary syndrome, osteoarthritis, rheumatoid arthritis, allergies, asthma, cancers, autoimmune diseases, pancreatitis, and cataract. Thus, Ndeprotection of Et 2-ethoxy-3-[4-(2-phthalimidoethoxy)phenyl]propionate with hydrazine hydrate in MeOH at room temperature for 1.5 h followed by amidation with 4-pyridin-2-ylbenzoic acid using carbonyl diimidazole in CH2Cl2 at room temperature for 1. 5 h followed by saponification with a mixture of 1 N aqueous NaOH and MeOH and acidification gave 3-[4-[2-(4-pyridin-2ylbenzoylamino)ethoxy]phenyl]propionic acid derivative (II.Na; R = H, R1 = Et) (III). III and (S)-II (R = H, R1 = 4-isopropoxyphenyl) in feed containing 0.01% at .apprx.10 mg drug/kg/day for 3 days lowered blood sugar level by 43 and 73%, resp. A capsule, a tablet, and a granule formulation containing III were prepared

L20 ANSWER 14 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1997:45114 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 126:148282

ORIGINAL REFERENCE NO.: 126:28607a,28610a

TITLE: Flavonol glycosides and phenolics from leaves of

Cordia dichotoma

AUTHOR(S): Wang, Yan; Ohtani, Kazuhiro; Kasai, Ryoji; Yamasaki,

Kazuo

CORPORATE SOURCE: Institute Pharmaceutical Sciences, Hiroshima

University School Medicine, Hiroshima, 734, Japan

SOURCE: Natural Medicines (1996), 50(5), 367

CODEN: NMEDEO; ISSN: 1340-3443

PUBLISHER: Japanese Society of Pharmacognosy

DOCUMENT TYPE: Journal LANGUAGE: English

AB Fresh leaves of Cordia dichotoma were extracted with MeOH. The concentrated extract was dissolved in water and successively extracted with hexane, EtOAc, and 1-BuOH. Six flavonol glycosides and two phenolic compds. were isolated from the butanol extract by a series of chromatog. Rosamarinic acid was the major constituent of the leaves of the plant, which may be responsible for the anti-inflammatory action of this plant.

TI Flavonol glycosides and phenolics from leaves of Cordia dichotoma REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

SO Natural Medicines (1996), 50(5), 367

CODEN: NMEDEO; ISSN: 1340-3443

IT 153-18-4P, Quercetin-3-O-rutinoside 604-80-8P,
Isorhamnetin-3-O-rutinoside 17297-56-2P 17650-84-9P,
Kaempferol-3-O-rutinoside 20283-92-5P, Rosmarinic acid
55696-57-6P 55804-74-5P 99353-00-1P, Methyl rosmarinate
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PUR (Purification or recovery); BIOL (Biological study); PREP (Preparation)

(flavonol glycosides and phenolics from leaves of Cordia dichotoma)

L20 ANSWER 15 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1996:515023 CAPLUS Full-text

DOCUMENT NUMBER: 125:212112

ORIGINAL REFERENCE NO.: 125:39394h,39395a

TITLE: 3α -Hydroxysteroid dehydrogenase inhibitory

actions of flavonoids and phenylpropanoids from

Schizonepeta spikes

AUTHOR(S): Matsuta, Muneto; Kanita, Rie; Saito, Yuji; Yamashita,

Akira

CORPORATE SOURCE: Kampo Res. Lab., Kanebo Ltd., Osaka, 534, Japan

SOURCE: Natural Medicines (1996), 50(3), 204-211

CODEN: NMEDEO; ISSN: 1340-3443

Japanese Society of Pharmacognosy

DOCUMENT TYPE: Journal LANGUAGE: Japanese

PUBLISHER:

AB Dry spikes of Schizonepeta tenuifolia Briquet (Labiatae) exhibited an inhibitory activity on 3α -hydroxysteroid dehydrogenase. From S. spikes, five flavonoids (hesperidin, hesperetin, apigenin, luteolin and ladanein), five phenylpropanoids (caffeic acid, rosmarinic acid, cinnamic acid, p-coumaric acid and 2,3-di-O-cinnamoyltartaric acid) and eight new caffeic acid derivs. (schizotenuins A-F) were isolated and their structures were elucidated on the basis of NMR, IR, UV, MS and other physicochem. evidences. The inhibitory activities of all of these compds. on 3α -hydroxysteroid dehydrogenase were stronger than that of aspirin. The pharmacol. effect of S. spikes was considered to be mainly due to luteolin, rosmarinic acid, schizotenuins A and C1, because they have strong inhibitory activities on 3α -hydroxysteroid dehydrogenase, and their contents were high.

ORIGINAL REFERENCE NO.: 121:14837a,14840a

TITLE: Competing O-H insertion and β -elimination in

rhodium carbenoid reactions; synthesis of

2-alkoxy-3-arylpropanoates

AUTHOR(S): Cox, Geoffrey G.; Haigh, David; Hindley, Richard M.;

Miller, David J.; Moody, Christopher J.

CORPORATE SOURCE: Dep. Chem., Loughborough Univ. Tech., Leicestershire,

KT18 5XO, UK

SOURCE: Tetrahedron Letters (1994), 35(19), 3139-42

CODEN: TELEAY; ISSN: 0040-4039

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 121:82641

GI

/ Structure 128 in file .gra /

AB Phodium(II) carboxylate catalyzed decomposition of diazo esters 3 (shown as I) and PhCH2C(CO2Et)N2 4 in the presence of alcs. or water results in formation of 2-alkoxy- or 2-hydroxy-3-arylpropanoates, resp., by O-H insertion in competition with cinnamates by elimination; the ratio of insertion to elimination is dramatically affected by the carboxylate ligand on rhodium. Use of methanol-d as the alc. confirms that the alkene does not arise by elimination from the initial alkoxyester product.

- TI Competing O-H insertion and β -elimination in rhodium carbenoid reactions; synthesis of 2-alkoxy-3-arylpropanoates
- TI Competing O-H insertion and β -elimination in rhodium carbenoid reactions; synthesis of 2-alkoxy-3-arylpropanoates
- SO Tetrahedron Letters (1994), 35(19), 3139-42 CODEN: TELEAY; ISSN: 0040-4039

AB Rhodium(II) carboxylate catalyzed decomposition of diazo esters 3 (shown as I) and PhCH2C(CO2Et)N2 4 in the presence of alcs. or water results in formation of 2-alkoxy- or 2-hydroxy-3-arylpropanoates, resp., by O-H insertion in competition with cinnamates by elimination; the ratio of insertion to elimination is dramatically affected by the carboxylate ligand on rhodium. Use of methanol-d as the alc. confirms that the alkene does not arise by elimination from the initial alkoxyester product.

L20 ANSWER 17 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1993:169109 CAPLUS Full-text

DOCUMENT NUMBER: 118:169109

ORIGINAL REFERENCE NO.: 118:29012h,29013a TITLE: Preparation of

(tetrazolylbiphenylmethyl)benzazepinones and related

compounds as growth hormone release promoters

INVENTOR(S): Fisher, Michael H.; Wyvratt, Matthew J.; Schoen,

William R.; Devita, Robert J.

PATENT ASSIGNEE(S): Merck and Co., Inc., USA SOURCE: PCT Int. Appl., 346 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PA:	TENT NO.			KIND	DATE	APPLICATION NO.	DATE
WO						WO 1992-US2271	19920319 <
						PL, RO, RU, SD	
	5206235					US 1992-839742	
EP	513974			A1	19921119	EP 1992-302143	19920312 <
EP	513974			B1	19960904		
	R: AT,	BE,	CH,	DE, D	K, ES, FR,	GB, GR, IT, LI, LU, NL,	PT, SE
AT	142206			T	19960915	AT 1992-302143	19920312 <
${ t IL}$	101206			A	19970218	IL 1992-101206	19920312 <
CA	2063185			A1	19920921	CA 1992-2063185	19920317 <
AU	9213012			A	19920924	AU 1992-13012	19920319 <
AU	653992			В2	19941020		
CN	1066070			А	19921111	CN 1992-102954	19920319 <
CN	1033584			С	19961218		
ZA	9202009			A	19921125	ZA 1992-2009	19920319 <
JP	06172316			A	19940621	JP 1992-112069	19920319 <
JP	08000814			В	19960110		
HU	66796			A2	19941228	ни 1992-915	19920319 <
RO	117326			В1	20020130		
US	5310737			A	19940510	US 1993-12190	19930202 <
	Y APPLN.		. :				19910320 <
							19920228 <
							19920319 <
OTHER SO	OURCE(S):			MARPA	T 118:1691		

GI

/ Structure 129 in file .gra /

AΒ Title compds. [I; L = (substituted) phenylene; n, w = 0, 1; p = 0-3; q = 0-4; X = CO, O, S, SO, SO2, CH(OH), CH:CH, imino; R1, R2, R7, R8 = H, halo, (perfluoro)alkyl, perfluoroalkoxy, cyano, NO2, (substituted) Ph, acyl(alkyl), etc.; R4, R5 = H, (substituted) Ph, alkyl, alkenyl, alkynyl, alkanoyloxy, alkoxycarbonyl, carboxy, CHO, amino; R4R5 = (CH2)rB(CH2)s; B = CH2, O, imino, S, SO, SO2; r, s = 1-3; R6 = H, alkyl, Ph, phenylalkyl; R9 = H, (substituted) tetrazolyl, acylalkyl, aminoalkyl, carbamoylalkyl, tetrazolylalkyl, tetrazolylphenyl, tetrazolylphenoxy, etc.; A = (CH2)xCR10R11(CH2)y; x, y = 0-3; R10, R11 = H, CF3, (substituted) alkyl, Ph, etc.; R10R11 = (CH2)t; t = 2-6; R10, R11 may be joined to R4 and/or R5], were prepared for promotion of release of growth hormone (no data). Thus, 3-benzyloxycarbonylamino-3-methyl-N-[2,3,4,5-tetrahydro-2-oxo-1H-1- benzazepin-3R-y1]butanamide (preparation from 3-azido-2,3,4,5-tetrahydro-1H-1-benzazepin-2-one given) was stirred 15 min with NaH in DMF; N-triphenylmethyl-5-[2-(4'-bromobiphen-4-yl)]tetrazole (preparation starting from 5-phenyl-2-trityltetrazole and 4-IC6H4Me given) in DMF was added and the mixture was stirred 90 min to give 95% coupling product, which was hydrogenated in MeOH over Pd(OH)2/C for 14 h to give 89% 3-amino-3methyl-N-[2,3,4,5-tetrahydro-2-oxo-1-[[2'-(1H-tetrazol-5-yl)][1,1'-biphenyl]-1-(2,3,4,5-tetrahydro-2-oxo-1-[[2'-(1H-tetrazol-5-yl)][1,1'-biphenyl]-1-(1H-tetrazol-5-yl)[1,1'4-y1]methy1]-1H-1-benzazepin-3R- y1]butanamide trifluoroacetate.

ACCESSION NUMBER: 1990:459705 CAPLUS Full-text

DOCUMENT NUMBER: 113:59705

ORIGINAL REFERENCE NO.: 113:10119a,10122a

Total syntheses of potassium lespedezate and potassium

isolespedezate, bioactive substances concerned with

circadian rhythm in nyctinastic plants

AUTHOR(S): Shigemori, Hideyuki; Miyoshi, Eiichi; Shizuri,

Yoshikazu; Yamamura, Shosuke

Fac. Sci. Technol., Keio Univ., Yokohama, 223, Japan CORPORATE SOURCE:

Tetrahedron Letters (1989), 30(46), 6389-92 SOURCE:

CODEN: TELEAY; ISSN: 0040-4039

DOCUMENT TYPE: Journal English LANGUAGE:

OTHER SOURCE(S): CASREACT 113:59705

GΙ

/ Structure 130 in file .gra /

AΒ Potassium lespedezate (I) and potassium isolespedezate (II) have been synthesized. The synthetic compds. have exhibited activities indistinguishable from the natural ones on leaf-opening of nyctinastic plants.

L20 ANSWER 19 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1975:443121 CAPLUS Full-text

DOCUMENT NUMBER: 83:43121

ORIGINAL REFERENCE NO.: 83:6811a,6814a

TITLE: Polyphenolic acids of Lithospermum ruderale (Boraginaceae). I. Isolation and structure

determination of lithospermic acid

Kelley, Charles J.; Mahajan, J. R.; Brooks, Lucille AUTHOR(S): C.; Neubert, Leonard A.; Breneman, W. R.; Carmack,

Marvin

CORPORATE SOURCE: Dep. Chem., Indiana Univ., Bloomington, IN, USA Journal of Organic Chemistry (1975), 40(12), SOURCE:

1804-15

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

GT For diagram(s), see printed CA Issue.

A structure is proposed for lithospermic acid (I), C27H22O12, the major polyphenolic acid of Lithospermum ruderale and several other plant species of the families, Boraginaceae and Labiatae. Chromatog. on Sephadex of aqueous exts. of the plant yields the di-K salt of I, together with salts of lesser constituents which include (R)-3-(3,4-dihydroxyphenyl)lactic acid, 2-(3,4dihydroxyphenyl)-3-carboxy-4-(2-carboxy-trans-vinyl)-7- hydroxycoumaran, and rosmarinic acid. Structures were deduced from spectral studies of the salts, the free acids, and also the methylated derivs., produced by the action of CH2N2 on the free acids or Me2SO4 on the salts.

L20 ANSWER 20 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN 1970:121115 CAPLUS Full-text ACCESSION NUMBER: DOCUMENT NUMBER: 72:121115

ORIGINAL REFERENCE NO.: 72:21759a,21762a

TITLE: Fukiic acid isolated from the hydrolysate of a

polyphenol in Petasites japonicus

AUTHOR(S): Sakamura, Sadao; Yoshihara, Teruhiko; Toyoda,

Katsuhiko

CORPORATE SOURCE: Dep. Agr. Chem., Hokkaido Univ., Sapporo, Japan

SOURCE: Agricultural and Biological Chemistry (1969

), 33(12), 1795-7

CODEN: ABCHA6; ISSN: 0002-1369

DOCUMENT TYPE: Journal LANGUAGE: English

AB A new polyphenol was isolated from leaves, leave stems, and flower stalks of P. japonicus, and named as fukinolic acid (I). Alkaline hydrolysis of I yielded equal moles of caffeic acid and a polyphenol , fukiic acid (II). II was crystallized as its monomethyl ester, 3,4- (HO)2C6H3CH2C(OH)(CO2H)C(OH)CO2Me, m. $188-90^{\circ}$. [α]2D2 40.5° (c 1, water), C12H14O8. II was also crystallized as its dimethyl ester, m. 140° , and the dimethyl ester of the dimethyl ether derivative m., $117-18^{\circ}$. The structure of II was suggested to be 3,4-(HO)2C6H3CH2C(OH)(CO2H)CH(OH)CO2H based on ir and mass spectra.

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L20 ANSWER 21 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1966:473090 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 65:73090

ORIGINAL REFERENCE NO.: 65:13587d-h,13588a

TITLE: Two diastereomeric forms of guaiacylglycerol

 β -(2-methoxyphenyl) ether and of gualacylglycerol

AUTHOR(S): Miksche, Gerhard E.; Gratzl, Josef; Fried-Matzka,

Maria

CORPORATE SOURCE: Chalmers Tekn. Hoegskol., Goteborg, Swed. SOURCE: Acta Chemica Scandinavica (1966), 20(4),

1038 - 43

CODEN: ACHSE7; ISSN: 0904-213X

DOCUMENT TYPE: Journal LANGUAGE: German

GI For diagram(s), see printed CA Issue.

AB The title compds. have been prepared as model substances to study lignin degradation products. Bromination of 109 g. Et benzylvanilloylacetate in 500 ml. CHC13 with 17.4 ml. Br in 100 ml. CHC13 in the presence of .apprx.75 g. (precipitated) CaCO3 gives 80% Et α -bromobenzylvanilloylacetate (I), 75-90° (decomposition), which is converted to 70% Et α -(2-methoxy-phenoxy) benzylvanilloylacetate (II), m. $87-9^{\circ}$, by treating with 60 g. K guaiacolate in 250 ml. HCONMe2 (DMF). Analogously, I with Na dihydroeugenol in DMF gives the 2-methoxy-4-propylphenoxy derivative which crystallizes in 2 modifications, a: m. 779° , b: m. $103-6^{\circ}$. I (5 q.) with 4.2 q. Na vanillate gives 4 q. 2-methoxy-4-formylphenoxy derivative, m. 118-20°. Reduction of 10 g. II in 150 ml. EtOH with H (1 g. Pd/C, 1 atmospheric) yields 75% Et erythro-2-(2-methoxyphenoxy)-3-hydroxy-3-(3-methoxy-4- hydroxyphenyl)propionate (III), m. 137-9° (EtOH-H2O); diacetyl derivative (from Ac2O-C5H5N) m. 106° (EtOH), b0.005 185-90°. Evaporation of the mother liquor gives 10% threo-form of III; diacetyl derivative, m. $98-100^{\circ}$ c. (EtOH). Reduction of 6 g. erythro-III in 100 ml. anhydrous tetrahydrofuran (THF) with 3 g. LiAlH4 in 200 ml. THF, 50° c., 6 hrs., gives, on addition of H2O, precipitation with dry-ice, and extraction with AcOEt, 3.8 g. erythro-quaiacylqlycerol- β -(2methoxyphenyl) ether (IV), m. 90-2°. This ether can also be obtained by reduction of II with LiAlH4. Analogously, reduction of 2.6 g. threo-III with LiAlH4 yields 93% 1.7 g.

threo-IV, m. 119-20°. Methylation of 2 g. III in 30 ml. 1:1 MeOH-dioxane with CH2N2 in Et2O yields Et erythro-2-(2-methoxyphenoxy)-3-hydroxy-3-(3,4-dimethoxyphenyl)propionate, m. 101-2°. The methylated threo derivative, m. 99-101°, is obtained analogously from threo-III. Methylation of IV gives 75% erythro-3,4-dimethoxyphenylglycerol- β -(2-methoxyphenyl) ether, m. 98-9°. The analogous methylated threo derivative obtained from threo-IV failed to crystallize; diacetate (from Ac2O-C5H5N) m. 96°. Treatment of 1 g. I in 10 ml. DMF with 2 g. anhydrous K acetate 1 hr. at 55° gives on dilution with H2O and extraction with Et2O 0.61 g. Et α -acetoxybenzylvanilloacetate, b0.201 185-8°. Further treatment (402 mg.)in 50 ml. EtOH with H over 0.1 g. Pd/C gives 307 mg. oily mixture of Et threo- and erythro-2-acetoxy-3-hydroxy-3-(3-methoxy-4-hydroxyphenyl)propionate. A crystalline triacetate (from Ac2O-C5H5N) of the erythro form is obtained, m. 110.5-11.5°, b0.007 145-50°, which can be reduced with LiAlH4 to yield the known erythro-guaiacylglycerol (V), m. 83-4° (Adler and Gustafsson, CA 59, 6301g).

L20 ANSWER 22 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1965:438715 CAPLUS Full-text

DOCUMENT NUMBER: 63:38715
ORIGINAL REFERENCE NO.: 63:6873c-g

TITLE: Synthesis of prephenic acid diethyl acetal and its

hydrolysis to phenylpyruvic acid and prephenic acid

AUTHOR(S): Plieninger, Hans; Arnold, Lothar; Fischer, Rolf;

Hoffmann, Werner

CORPORATE SOURCE: Univ. Heidelberg, Germany

SOURCE: Chemische Berichte (1965), 98(6), 1774-81

CODEN: CHBEAM; ISSN: 0009-2940

DOCUMENT TYPE: Journal LANGUAGE: German

GI For diagram(s), see printed CA Issue.

AΒ I and the di-Et acetal (II) of III were prepared The time at which the maximum yield of III can be obtained by the acid hydrolysis of II was calculated from kinetic data and the calcn. confirmed by the experiment; III was formed together with larger amts. of PhCH2COCO2H (IV). Di-Et 2cyclohexen-4-one-1-carboxylate-1-pyruvate di-Et acetal (35.6 g.), b0.1 155°, n25D 1.4087, in 350 cc. tert-BuOH refluxed 5 hrs. with stirring with 5 cc. AcOH and 11 g. SeO2, treated with an addnl. 11 g. SeO2, and again refluxed 5 hrs., and the product shaken in Et2O 6 hrs. with 10 g. deactivated Raney Ni yielded 25 g. I, b0.1 150°, n20D 1.4840. I (35.6 mg.), 5.0 cc. EtOH, and 4.0cc. 0.1N NaOH heated 0.5 hr. at 50° , treated with 5 cc. N HCl, and heated 15 min. at 50° , and the mixture cooled and diluted with H2O to 100 cc. gave a solution containing 18 mg. p-HOC6H4CH2COCO2H. I (1.0 g.), 100 mg. Na, and 5 cc. EtOH heated 3 hrs. at 50° gave 0.70 g. p-HOC6H4CH24C(OEt)2CO2Et (IV). I (7.1 g.) in 10 cc. EtOH added dropwise with stirring during 15 min. at 5° to 0.40 g. NaBH4 in 40 cc. EtOH and kept 15 min. at 20° yielded 5.3 g. oily di-Et ester (V) of II, n25D 1.4665, and 1.1 g. IV. V (3.6 g.) in 30 cc. EtOH treated 2 days at room temperature with 1.6 g. NaOH in 20 cc. H2O and evaporated to 15 cc., buffered to pH 8 with N HCl, and diluted with H2O to 25 cc. gave a 0.4M solution; a 0.1-cc. portion and 4 cc. N HCl heated 15 min. at 50° and the mixture adjusted to pH 14 with N NaOH gave a 0.3M II solution; a 6.25-cc. portion stirred with 4.25 g. Ba(OAc)2, centrifuged to remove some solid, diluted with about 200 cc. EtOH, and kept several hrs. yielded 1.3 q. Ba salt; a 0.5-g. portion in 10 cc. H2O hydrogenated over PdBaSO4, filtered, and treated 20 hrs. with 2,4-(O2N)2C6H3NHNH2 in 2N HCl, and the precipitate chromatographed on paper showed the presence of the 2,4dinitrophenylhydrazones of cis- and trans-tetrahydroprephenic acid (relative to OH and CO2H groups). II (0.3M aqueous solution) (10 cc.) adjusted with N HCl to pH 1.8, kept 10 min. at 20°, neutralized with N NaOH, and treated with

1.0 g. Ba(OAc)2 in 5 cc, H2O gave 0.5 g. Ba salt of a 6:50 mole ratio mixture of III and IV.

L20 ANSWER 23 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1965:30781 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 62:30781
ORIGINAL REFERENCE NO.: 62:5491f-g

TITLE: The nonvolatile acids of succulent plants exhibiting a

marked diurnal oscillation in their acid content. I. The detection of piscidic acid in Agave americana

AUTHOR(S): Nordal, Arnold; Ogner, Gunnar

CORPORATE SOURCE: Univ. Oslo, Norway

SOURCE: Acta Chemica Scandinavica (1964), 18(8),

1979-83

CODEN: ACHSE7; ISSN: 0904-213X

DOCUMENT TYPE: Journal LANGUAGE: English

AB Piscidic acid (I) was detected in the leaves of A. americana by the following procedure: the acid mixture, isolated over the Pb salts from fresh leaves of the plant, was converted to the corresponding Me and Et esters, and the ester mixts. were fractionated in vacuo. From the individual fractions the hydrazides and benzylidene hydrazides were prepared and examined A crystalline hydrazide (C11H16O5N4; m. 185-187°), corresponding to the hydrazide of I, was isolated from the highest boiling fractions, and from this hydrazide the corresponding benzylidene hydrazide (C25H25O5N4; m. 136-138°) was prepared From the highest boiling fractions of the Me ester mixture, a crystalline ester (C13H16O7; m. 126-127°), corresponding to the Me ester of I, was isolated and from this an acetyl derivative (m. 84°) was prepared The properties of these 4 derivs. and the ir spectra of the ester identified the acid in question as I.

- TI The nonvolatile acids of succulent plants exhibiting a marked diurnal oscillation in their acid content. I. The detection of piscidic acid in Agave americana
- SO Acta Chemica Scandinavica (1964), 18(8), 1979-83 CODEN: ACHSE7; ISSN: 0904-213X
- AB Piscidic acid (I) was detected in the leaves of A. americana by the following procedure: the acid mixture, isolated over the Pb salts from fresh leaves of the plant, was converted to the corresponding Me and Et esters, and the ester mixts. were fractionated in vacuo. From the individual fractions the hydrazides and benzylidene hydrazides were prepared and examined A crystalline hydrazide (C11H16O5N4; m. 185-187°), corresponding to the hydrazide of I, was isolated from the highest boiling fractions, and from this hydrazide the corresponding benzylidene hydrazide (C25H25O5N4; m. 136-138°) was prepared From the highest boiling fractions of the Me ester mixture, a crystalline ester (C13H16O7; m. 126-127°), corresponding to the Me ester of I, was isolated and from this an acetyl derivative (m. 84°) was prepared The properties of these 4 derivs. and the ir spectra of the ester identified the acid in question as I.

L20 ANSWER 24 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1963:435348 CAPLUS Full-text

DOCUMENT NUMBER: 59:35348

ORIGINAL REFERENCE NO.: 59:6301g-h,6302a-d

TITLE: Preparation of the threo- and erythro-forms of DL-quaiacylqlycerol and of DL-veratrylqlycerol

AUTHOR(S): Adler, Erich; Gustafsson, Bo

CORPORATE SOURCE: Chalmers Tek. Hogskola, Goteborg, Swed. SOURCE: Acta Chemica Scandinavica (1963), 17, 27-36

CODEN: ACHSE7; ISSN: 0904-213X

DOCUMENT TYPE: Journal LANGUAGE: German

OTHER SOURCE(S): CASREACT 59:35348

cf. CA 48, 5147g. Hydrogenation of 3,4-dimethoxyphenylpropiolic acid in EtOH AB with Lindlar catalyst 50 min. gave 70% cis-methylferulic acid (I), m. 104°, which with CH2N2 yielded 90% Me ester, prisms, m. 92-3°. Reduction of 30 g. Me trans-methylferulate (II) in 200 cc. dioxane with 5.7 g. LiAlH4 in 500 cc. absolute Et20, addition of H2O and dilute H2SO4, extraction with CHCl8, and distillation of the residue of the CHC13 extract gave 70% methylconiferyl alc., b5 110-20°, needles, m. 79-80°; Ac derivative (III) b12 190-5°. Treating 80 mg. III in 3 cc. Et20-C5H5N (25:1) 16 hrs. with 0.1 g. 0s04 in 2 cc. Et20 and hydrolyzing the osmic ester in 2 cc. Et0H with 0.65 g. Na2SO3 in $3 \, \text{cc. H2O} \, 1 \, \text{hr. at} \, 100^{\circ}$, evaporating the filtered solution in vacuo, and extracting the residue with CHCl3 gave 85% DL-threoveratrylglycerol (IV), m. 109-10°. Treating II similarly with OsO4 and acetylating the hydrolyzed Os ester gave 60% DL-threo- α , β -diacetoxymethylhydroferulic acid (V), prisms, m. 148-9°. Esterification of V with CH2N2 followed by reduction yielded IV. Refluxing 15 g. Me α -bromo- β -acetoxymethyl-hydroferulate in 65 cc. AcOH and 65 cc. Ac2O 45 min. with 7 g. AgOAc, treating the filtered solution with H2O, evaporating the solution in vacuo, extracting the residue with CHC13, evaporating the washed (NaHCO3) solution, and crystallizing the residue from Et20-C6H14 gave 41% Me threo- α , β -diacetoxymethylhydroferulate (VI), m. 102-3°. Concentration of the mother liquor gave 40% erythro isomer (VII), prisms, m. 71-3°. VI was also obtained in 10% yield when 4.44 g. II was oxidized with KMnO4 at -50° according to Riiber (CA 9, 2244). Reduction of VII with LiAlH4 gave DL-erythroveratrylglycerol (VIII), plates, m. 92-3°, λ maximum 278 m μ (log ϵ 3.52). The infrared spectra of IV and VIII differed distinctly. When Me erythro-diacetoxyacetylhydroferulate was reduced with LiAlH4 (cf. loc. cit.) and the acid step was avoided by neutralizing the reaction mixture with AcOH, 55% DL-erythro-guaiacylglycerol (IX), m. $83-4^{\circ}$ was obtained; IX tetraacetate m. 86-8°. IX and CH2N2 gave 90% VIII. Treating 0.31 g. IX with 0.1N H2SO4 neutralizing the mixture with BaCO8, evaporating the filtered solution in vacuo, and treating the residue with moist EtOAc gave 0.18 g. unchanged IX and, from the mother liquor, 0.05 q. threo-DL-quaiacylqlycerol (X); tetraacetate m. 113-14°. Benzyl-coniferyl alc. benzoate (0.515 g.) (Freudenberg and Achtzehn, CA 50, 1661h) was treated in 12 cc. Et20-C5H5N with 0.35 g. OsO4 and the precipitate formed was boiled 1 hr. with 2.3 g. Na2SO3 in 10 cc. H2O, giving threo-(O-benzylquaiacyl)glycerol, m. 101°, which when treated in EtOH with prehydrogenated PdCl2-BaSO4, yielded X as a sirup; tetraacetate, 70%, m. 113-14°. X and CH2N2 gave 70% IV. Careful fractionation of the mother liquor of the Me erythro- α , β diacetoxyacetylhydroferulate gave 10% threo isomer, prisms, m. 81-2°, which with LiAlH4 gave 55% X. Treating 0.074 g. trans-methylisoeugenol (XI) with OsO4 gave 75% DL-threo-methylisoeugenol glycol, m. 90°, which was also obtained in 15% yield when 1.93 g. XI was treated at -50° with 1.96 g. KMnO4. Acetylation of 3.3 g. XI in 3 cc. AcOH with 6.6 g. Pb(OAc) 4 and reduction of the α , β -diacetoxyveratrylpropane formed with LiAlH4 gave 2 g. of a product, m. 80-100°, which on fractional crystallization yielded crythro-methylisoeugenol glycol, m. 123°, and the threo isomer, m. 88°.

L20 ANSWER 25 OF 26 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1957:5337 CAPLUS Full-text

DOCUMENT NUMBER: 51:5337

ORIGINAL REFERENCE NO.: 51:1084e-i,1085a-d

TITLE: Synthesis and reactions of quaiacylglycerol

AUTHOR(S): Stumpf, Walter; Rumpf, Gunther CORPORATE SOURCE: Univ. Heidelberg, Germany

SOURCE: Annalen der Chemie, Justus Liebigs (1956),

599, 51-60 CODEN: 9X224Y

DOCUMENT TYPE: Journal LANGUAGE: Unavailable OTHER SOURCE(S): CASREACT 51:5337

3,4-MeO(PhCH2O)C6H3CHO (138 g.) in 230 cc. AcOMe, after standing overnight with 17.1 g. Na wire and 2 cc. MeOH was triturated carefully with 230 cc. AcOMe, kept another 48 hrs. at room temperature, refluxed 1 hr. with 230 cc. absolute Et20, and shaken with 740 cc. H2SO4. The organic phase washed with aqueous NaHCO3 and H2O, dried and evaporated gave 95.5% crude and 83% pure 3,4-MeO(PhCH2O)C6H3CH:CHCO2Me (I), m. 98-9° (from MeOH or PrOH). To 7.16 g. I in 24 cc. CHCl2 at 0°-5° were added dropwise (over a 1.5-hr. period) 3.84 g. Br; the mixture after 1 hr. at 0° was evaporated giving the crude di-Br derivative (II) of I (not weighed or analyzed), 22.9 g. of which were added to 15 g. dry AcOK in 150 cc. AcOH and 50 cc. Ac2O, heated 9 hrs. on a steam bath, then boiled 8 hrs., filtered, and concentrated in vacuo to incipient crystallization, treated with H2O and extracted with ET2O; the washed and dried extract evaporated in vacuo gave 19 g. sirup, a small sample of which, triturated with MeOH, gave seed crystals of (III), C22H24O8, m. sharply 112.5-13.5° (after 2 crystns. from MeOH). The main portion of the sirup, inoculated with III, crystallized gradually giving 8.25 g. of what was probably a mixture of isomeric forms of 3,4-MeO(PhCH2O)C6H3CH(OAc)CH(OAc)CO2Me (IV), m. poorly $90-100^{\circ}$ (even after repeated crystallization from MeOH). In another experiment in which 169 g. crude II was heated 15 hrs. at 100° with 110 g. AcOK, 800 cc. AcOH, and 400 cc. Ac2O, a red sirup was formed, which, with III, gave 52.45 g. IV, leaflets, m. poorly 89-98°, the mother liquors from which gave I. A series of fully described attempts were made to fractionate IV into its component (racemic) isomers, but although 3 fractions were obtained, m., resp., $85.5-87^{\circ}$, $86-91^{\circ}$, and $88-91^{\circ}$, none of these was homogeneous. IV (m. $89-98^{\circ}$)(12.5 g.) in 50 cc. dry AcOMe was hydrogenated with 0.75 g. 2% pd-BaSO4. After 1.5 hrs. 745 cc. H had been taken up. The filtered, evaporated solution gave 3,4-MeO(HO) C6H3CH(OAc) CH(OAc) CO2Me (V), viscous, uncrystallizable pale yellow sirup. V (8.34 g.) in 100 cc. absolute Et2O was added dropwise to 5.6 g. LiAlH4 in 200 cc. Et2O, and after 5 hrs. at room temperature was refluxed 2 hrs., cooled to 0° in a stream of CO2 and treated dropwise with H2O, shaken with H2O saturated with CO2, the Et2O layer separated and the aqueous phase extracted continuously for 7 days with peroxide-free Et20 in a Perforator, using fresh Et20 after 28 hrs. (when a sirup separated from the Et2O-phase). The various combined Et2O exts. evaporated in vacuo gave 2.7 g. crude resinous guaiacylglycerol (VI), which was boiled briefly with 200 cc. H2O, filtered and reextd. twice with Et2O. The aqueous phase (in which VI is very soluble) was evaporated to dryness in vacuo under N, giving 1.98 g. (36.3%) purified VI, C10H14O5, yellow sirup (after drying 14 hrs. at 34° in vacuo and 2 days at 20° over P2O5). VI is difficultly soluble in Et20 and C6H6 and could not be crystallized IV (m. 87-93°) (21.2 g.) in 750 cc. absolute Et20 was stirred into a mixture of 10 g. LiAlH4 and 200 cc. Et20, and treated as in the case of V. The resultant aqueous solution was filtered and extracted 40 hrs. with Et20; the Et20 layer yielded 2.15 g. PhCH2 derivative (VII) of VI, m. 68-74° (from C6H6), m. 99-100.5° (from MeOH by addition of Et2O and petr. ether to incipient cloudiness, or from AcOEt). Even this purified VII may be a mixture of racemic isomers. VI (1.81 g.) and Na2S2O5 in 85 cc. H2O was heated and shaken 19 hrs. at 135° in a sealed tube, then freed from SO2 and extracted 40 hrs. with Et20 giving 0.15 g. impure 4,3-HO(MeO)C6H3CH(SO3H)CH(OH)CH2OH, 81% of which was soluble in cold H2O forming a pale pink Ba salt (VIII) (containing 32.68% C and 21.3% Ba; calculated 34.71 and 19.85%, resp.). Oxidations with NaIO4 were carried out with various phenolic compds. or their derivs., and results are given in terms of moles NaIO4 consumed per mole of compound within a specific time period. No NaIO4 was consumed by veratrole within 17.5 hrs. The consumption of NaIO4 by

PhOH, vanillin, and p-cresol was very slight (0.14-0.33 mole within 21.5-24 hrs.). V, VI, guaiacol, cresol, guaiacylethylcarbinol, VIII, 1,4-C6H4(OH)2, catechol, and pyrogallol all consumed appreciable amts. of NaIO4 within relatively short periods. On oxidation all guaiacyl compds. gave red solns., the color being ascribed to quinoid oxidation products. Oxidation data are discussed at length.

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L16
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SEARCH TIME: 00.00.01
L17
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L17 232 ANSWERS
                  REGISTRY COPYRIGHT 2009 ACS on STN
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L27 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                       2008:994060 CAPLUS Full-text
DOCUMENT NUMBER:
                        149:306485
                        Di-m-Chlorobis[Bis-(cyclooctene)rhodium]
TITLE:
                        Judd, Andrew S.
AUTHOR(S):
CORPORATE SOURCE:
                        USA
                        e-EROS Encyclopedia of Reagents for Organic Synthesis
SOURCE:
                        (2001), No pp. given. John Wiley & Sons,
                        Ltd.: Chichester, UK.
                        CODEN: 69KUHI
                        URL: http://www3.interscience.wiley.com/cgi-
                        bin/mrwhome/104554785/HOME
DOCUMENT TYPE:
                        Conference; General Review; (online computer file)
LANGUAGE:
                        English
OTHER SOURCE(S):
                        CASREACT 149:306485
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AB
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ΤI
SO
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L27 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                        2004:2837 CAPLUS Full-text
DOCUMENT NUMBER:
                        140:59411
                        Preparation of phenoxyalkanamides as amide linker
TITLE:
                        peroxisome proliferator activated receptor agonists
                        for treating and/or preventing diabetes mellitus and
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syndrome X

INVENTOR(S): Ferritto Crespo, Rafael; Martin, Jose Alfredo;

Martin-Ortega, Finger Maria Dolores; Rojo Garcia, Isabel; Shen, Quanrong; Warshawsky, Alan M.; Xu,

Yanping

PATENT ASSIGNEE(S): Eli Lilly and Company, USA SOURCE: PCT Int. Appl., 168 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	CENT N	10.			KINI)	DATE			APPI	LICAT	ION :	NO.		D.	ATE		
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AΒ The present invention is directed to phenoxyalkanamides (shown as I; variables defined below; e.g. II), compns., and their use as peroxisome proliferator activated receptor agonists for treating and/or preventing diabetes mellitus and syndrome X. The binding and cotransfection efficacy values found for compds. of this invention that are useful for modulating a PPARlpha receptor are about <100 nM and >50%, resp. Although the methods of preparation are not claimed, .apprx.140 example prepns. of I are included. For example, II was prepared in 3 steps starting from (2S)-2-ethoxy-3-(4-hydroxyphenyl)propionic acid Me ester, (2S)-2-hydroxypropionic acid benzyl ester and involving intermediates (2S)-3-[4-[[(1R)-1-[(benzyloxy)carbonyl]ethyl]oxy]phenyl]-2ethoxypropionic acid Et ester and (2S)-3-[4-[((1R)-1carboxyethyl)oxy]phenyl]-2-ethoxypropionic acid. For I: R1 = H, C1-C8 alkyl,

C3-C6 cycloalkyl, aryl-C0-4-alkyl, heteroaryl-C0-4-alkyl, aminoC1-C4alkyl, C3-C6 cycloalkylaryl-C0-2-alkyl, arylheteroC1-C8alkyl, -CHC(0)C1-C4 alkoxy, C0-4-alkyl-C(0)heteroC1-C8alkyl, and -CH2C(0)-R15R16. R2 = C1-C8 alkyl, C3-C6 cycloalkyl, aryl-C0-C4-alkyl, heteroaryl-C0-C4-alkyl, heteroC1-C6cycloalkylaryl, heteroC1-C6cycloalkylarylC1-C4alkyl, aminoC1-C4alkyl, C3-C6 cycloalkylaryl-C0-C2-alkyl, arylheteroC1-C8alkyl, C0-C4-alkyl-C(0)heteroC1-C8alkyl, -CH(C(0)OCH3)benzyl, and -CH2C(0)R15''R16''. R1 and R2 together may form a heterocyclic ring which heterocyclic ring is (un)substituted with 1-3 substituents R1' and which heterocyclic ring is optionally fused with an aryl; E = C(R3)(R4)A, (CH2)nCOOR13, aryl-C0-C4-alkyl, thio-C1-C4-alkyl, thioaryl, aryl-C1-C4alkoxy, C1-C4alkoxy C1-C4alkyl, aminoaryl, and aminoC1-C4alkyl, R5 and R6 = H, C1-C8 alkyl, aryl-C0-C4-alkyl, heteroaryl-C0-C4-alkyl, and -CH2C(0)R17R18.

L27 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2001:416876 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 135:33368
TITLE: Preparation of

(S)-2-ethoxy-3-(4-hydroxyphenyl) propanoates by

resolution using chiral amines.

INVENTOR(S): Andersson, Kjell; Fischer, Alan Eric; Ioannidis,

Panagiotis; Larsson, Magnus; Larsson, Maria;

Sivadasan, Sivaprasad Astrazeneca AB, Swed.

PATENT ASSIGNEE(S): Astrazeneca AB, Swed. SOURCE: PCT Int. Appl., 28 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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				WO	2000-SE2382	\mathbb{W}	20001129	<
PRIORITY	APPLN. INFO.:			SE	1999-4415	Α	19991203	<
US	20060069283	A1	20060330	US	2005-282247		20051118	<
US	7002037	В2	20060221					
US	20030139474	A1	20030724	US	2002-148818		20021113	<
KR	794091	B1	20080110	KR	2002-707070		20020601	<
ИО	2002002604	A	20020711	ИО	2002-2604		20020531	<
MX	2002PA05321	A	20021206	MX	2002-PA5321		20020529	<

OTHER SOURCE(S): CASREACT 135:33368; MARPAT 135:33368

GΙ

/ Structure 181 in file .gra /

AΒ Title compds. (I; R2 = H, protecting group; the aryl ring may be halogenated) (II; Q = H, protecting group) with chiral amines to form the diastereomeric salts, which were separated by crystallization followed by removal of the amine and optional deprotection and protection steps. Thus, 2-ethoxy-3-(4methoxyphenyl) propionic acid (preparation given) in iPrOAc at 0-5° was treated with (S)-1-(1-naphthyl) ethylamine followed by heating to $75-80^{\circ}$, cooling, and seeding with (S)-2-ethoxy-3-(4-methoxyphenyl) propionic acid (S)-1-(1-methoxyphenyl)naphthyl)ethylamine salt (III) to precipitate crude III, which was recrystd. from iPrOAc to give 74% III. III in PhMe was treated with aqueous NaOH and then with aqueous HCl followed by extraction of the aqueous layer with EtOAc to give an EtOAc solution of (S)-2-ethoxy-3-(4-methoxyphenyl) propionic acid. The EtOAc was replaced with N-methylpyrrolidone and the resulting solution was heated with NaOH and octanethiol at 115-125° to give 52% (S)-2-ethoxy-3-(4hydroxyphenyl)propionic acid in 99.8% chemical purity and 97.8% enantiomeric excess.

L27 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1998:82175 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 128:167302

ORIGINAL REFERENCE NO.: 128:32973a,32976a

TITLE: A non-enzymic synthesis of (S)-(-)-rosmarinic acid and

a study of a biomimetic route to (+)-rabdosiin

AUTHOR(S): Boqucki, David E.; Charlton, James L.

CORPORATE SOURCE: Dep. Chem., Univ. Manitoba, Winnipeg, MB, R3T 2N2,

Can.

SOURCE: Canadian Journal of Chemistry (1997),

75(12), 1783-1794

CODEN: CJCHAG; ISSN: 0008-4042

PUBLISHER: National Research Council of Canada

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 128:167302

GΙ

 $^{^{\}star}$ STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The synthesis of (S)-(-)-rosmarinic acid (I; R = R1 = H) in 9% overall yield is described. The synthesis was achieved by a convergent route in which 3-(3',4'-dihydroxyphenyl)-(S)-lactic acid and caffeic acid, (E)-3,4-(HO)2C6H3CH:CHCO2H, both appropriately protected, were coupled to produce a pentaallyl precursor I (R = R1 = CH2CH:CH2), which was then deprotected to give (S)-(-)-rosmarinic acid (I; R = R1 = H). A triallyl derivative I (R = H, R1 = CH2CH:CH2) was similarly prepared and converted to (+)-rabdosiin (II) and its (1R,2S) isomer via a biomimetic oxidative free radical coupling-cyclization followed by deallylation. The coupling-cyclization gave a ratio of rabdosiin diastereomers unlike that found in nature. A preliminary study showed that Me (R)-mandelyl sinapate (III) could be dimerized diastereoselectively to give a 1,2-trans thomasidioate diester (IV).

L27 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1996:436105 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 125:109268

ORIGINAL REFERENCE NO.: 125:20363a,20366a

TITLE: Separation of rosmarinic acid enantiomers by three

different chromatographic methods (HPLC, CE, GC) and the determination of rosmarinic acid in Hedera helix L

AUTHOR(S): Trute, Andreas; Nahrstedt, Adolf

CORPORATE SOURCE: Inst. Pharm. Biol. Phytochem., Wilhelms-Univ.,

Muenster, D-48149, Germany

SOURCE: Phytochemical Analysis (1996), 7(4), 204-208

CODEN: PHANEL; ISSN: 0958-0344

PUBLISHER: Wiley
DOCUMENT TYPE: Journal
LANGUAGE: English

- AB Three methods with HPLC, capillary electrophoresis (CE), and gas chromatog. (GC) were developed for the separation of enantiomers of rosmarinic acid (RA). Chiral resolution of underivatized RA was achieved on a chiral AGP HPLC column and by micellar electrokinetic capillary chromatog. Alternatively, after derivatization, the resulting diastereomeric 3,4-dihydroxyphenyllactic acid(-)-menthyl esters were separated by non-chiral GC. These methods allow reliable and rapid determination of the enantiomeric ratio of RA samples. RA obtained from Hedera helix L. (Araliaceae) was determined to be (R)-(+)-RA. In contrast to published data, the optical rotation value of rosmarinic acid was + 106°.
- TI Separation of rosmarinic acid enantiomers by three different chromatographic methods (HPLC, CE, GC) and the determination of rosmarinic acid in Hedera helix ${\tt L}$
- SO Phytochemical Analysis (1996), 7(4), 204-208 CODEN: PHANEL; ISSN: 0958-0344
- Three methods with HPLC, capillary electrophoresis (CE), and gas chromatog. (GC) were developed for the separation of enantiomers of rosmarinic acid (RA). Chiral resolution of underivatized RA was achieved on a chiral AGP HPLC column and by micellar electrokinetic capillary chromatog. Alternatively, after derivatization, the resulting diastereomeric 3,4-dihydroxyphenyllactic acid(-)-menthyl esters were separated by non-chiral GC. These methods allow reliable and rapid determination of the enantiomeric ratio of RA samples. RA obtained from Hedera helix L. (Araliaceae) was determined to be (R)-(+)-RA. In contrast to published data, the optical rotation value of rosmarinic acid was + 106°.

DOCUMENT NUMBER: 115:90598

ORIGINAL REFERENCE NO.: 115:15571a,15574a

TITLE: Chemo-enzymic synthesis of rosmarinic acid

AUTHOR(S): Pabsch, K.; Petersen, M.; Rao, N. N.; Alfermann, A.

W.; Wandrey, C.

CORPORATE SOURCE: Inst. Biotechnol., Res. Cent. Juelich, Juelich,

D-5170, Germany

SOURCE: Recueil des Travaux Chimiques des Pays-Bas (

1991), 110(05), 199-205

CODEN: RTCPA3; ISSN: 0165-0513

DOCUMENT TYPE: Journal LANGUAGE: English

GΙ

/ Structure 182 in file .gra /

AB (R)-(+)-3-(3,4-Dihydroxyphenyl)lactic acid (I) was produced in an enzyme membrane reactor with an optical purity >99%. Complete reactor modeling of the process was done and the theor. evaluated data was verified by a continuous experiment with volumetric yields of >1000 g/L-day. For the synthesis of CoA-activated caffeic acid, a new method was found to prepare the ester via the caffeic acid imidazolide. In a batch experiment, enzymically prepared I and chemical synthesized CoA caffeic ester were converted to rosmarinic acid (II) by an enzyme isolated from Coleus blumei cell cultures.

L27 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1988:492624 CAPLUS Full-text

DOCUMENT NUMBER: 109:92624

ORIGINAL REFERENCE NO.: 109:15441a,15444a

TITLE: Stereostructure of salvianolic acid B and isolation of

salvianolic acid C from Salvia miltiorrhiza

AUTHOR(S): Ai, Chunbo; Li, Lianniang

CORPORATE SOURCE: Inst. Mater. Med., Chin. Acad. Med. Sci., Beijing,

Peop. Rep. China

SOURCE: Journal of Natural Products (1988), 51(1),

145 - 9

CODEN: JNPRDF; ISSN: 0163-3864

DOCUMENT TYPE: Journal LANGUAGE: English

GΙ

/ Structure 183 in file .gra /

The depside salvianolic acid B (I), isolated from roots of S. miltiorrhiza was assigned a 2R,3R configuration and a 2β -pseudoequatorial aryl, 3α -pseudoaxial carboxyl conformation based on chemical degradation and spectral anal. Salvianolic acid C (II) was isolated from S. miltiorrhiza roots and the structure determined. The fact that salvinolic acid A was converted to II in a TLC plate impregnated with 2% formic acid suggests II to be the cyclization product of the former.

L27 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1975:443121 CAPLUS Full-text

DOCUMENT NUMBER: 83:43121

ORIGINAL REFERENCE NO.: 83:6811a,6814a

TITLE: Polyphenolic acids of Lithospermum ruderale (Boraginaceae). I. Isolation and structure

determination of lithospermic acid

AUTHOR(S): Kelley, Charles J.; Mahajan, J. R.; Brooks, Lucille C.; Neubert, Leonard A.; Breneman, W. R.; Carmack,

Marvin

CORPORATE SOURCE: Dep. Chem., Indiana Univ., Bloomington, IN, USA

SOURCE: Journal of Organic Chemistry (1975), 40(12),

1804 - 15

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

GI For diagram(s), see printed CA Issue.

AB A structure is proposed for lithospermic acid (I), C27H22O12, the major polyphenolic acid of Lithospermum ruderale and several other plant species of the families, Boraginaceae and Labiatae. Chromatog. on Sephadex of aqueous exts. of the plant yields the di-K salt of I, together with salts of lesser constituents which include (R)-3-(3,4-dihydroxyphenyl)lactic acid, 2-(3,4-dihydroxyphenyl)-3-carboxy-4-(2-carboxy-trans-vinyl)-7- hydroxycoumaran, and rosmarinic acid. Structures were deduced from spectral studies of the salts, the free acids, and also the methylated derivs., produced by the action of CH2N2 on the free acids or Me2SO4 on the salts.

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=> s 117 and 'asymmetric hydrogenation'
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2028 L17

76842 'ASYMMETRIC'

31 'ASYMMETRICS'

76873 'ASYMMETRIC'

('ASYMMETRIC' OR 'ASYMMETRICS')

147845 'ASYM'

6 'ASYMS'

147848 'ASYM'

('ASYM' OR 'ASYMS')

171543 'ASYMMETRIC'

('ASYMMETRIC' OR 'ASYM')

184239 'HYDROGENATION'

2478 'HYDROGENATIONS'

184500 'HYDROGENATION'

('HYDROGENATION' OR 'HYDROGENATIONS')

4062 'ASYMMETRIC HYDROGENATION'

('ASYMMETRIC'(W)'HYDROGENATION')

L28 2 L17 AND 'ASYMMETRIC HYDROGENATION'

=> d 128 ibib abs 1-2

L28 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2007:696717 CAPLUS Full-text

DOCUMENT NUMBER: 147:95305

TITLE: Process for the preparation of enantiomer-enriched

2-alkoxy-3-arylpropionic acids by asymmetric hydrogenation of substituted 2-alkoxycinnamic

acids

INVENTOR(S): Woltering, Michael; Bunlaksananusorn, Tanasri;

Gerlach, Arne

PATENT ASSIGNEE(S): Saltigo G.m.b.H., Germany SOURCE: Eur. Pat. Appl., 16pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
EP 1801093	A1 20070627		EP 2006-25546	20061211	
R: AT, BE, BG,	CH, CY	, CZ, DE,	DK, EE, ES, FI, FR, GE	B, GR, HU, IE,	
IS, IT, LI,	LT, LU	, LV, MC,	NL, PL, PT, RO, SE, SI	[, SK, TR, AL,	
BA, HR, MK,	YU				
DE 102005061472	A1	20070705	DE 2005-10200506147	72 20051222	
US 20070149804	A1	20070628	US 2006-635302	20061207	
US 7429676	B2	20080930			
CN 1986516	A	20070627	CN 2006-10168677	20061222	
PRIORITY APPLN. INFO.:			DE 2005-10200506147	72A 20051222	

CASREACT 147:95305; MARPAT 147:95305 OTHER SOURCE(S):

Chiral 2-alkoxy-3-arylpropanoic acids R2nC6H5-nCH2CH(OR1)CO2H or their alkali metal salts [1; R1 = (un)substituted C1-18 alkyl, C4-24 aryl, C5-18 arylalkyl; R2 = OH, halo, (alkyl)amino, C1-18 alkyl(oxy), C4-24 aryl, C5-18 arylalkyl, C1-18 alkylsulfonyl(amino), acyl(amino), acyloxy, preferably R2 = OH; n = 1-5, preferably n = 1], useful as peroxisome proliferator activated receptors (PPAR) agonists, were prepared by an improved process comprising transition metal-catalyzed asym. hydrogenation of the corresponding cinnamic acids R2nC6H5-nCH:C(OR1)CO2H (2; same R, n) in the presence of at least one protic solvent. Compds. 2 were preferably prepared by Perkin condensation of benzaldehydes R2nC6H5-nCHO (3; same R2, n) with 2-alkoxyacetates R10CH2CO2R3 (4; same R1; R3 = H, C1-18 alkyl, preferably C1-6 alkyl). In an example, sodium 4-hydroxy- α -methoxybenzenepropanoate (α S)-4-HOC6H4CH2CH(OMe)CO2Na was prepared in 53% yield and 92% ee by asym. hydrogenation of 200.0 mmol of (2Z)-4-HOC6H4CH:C(OMe)CO2H catalyzed by 0.5 mmol of [Ir(COD)Cl]2 and 1.0 mmol of (S,S)-2,4-bis(diphenylphosphino)pentane in 240 mL of iso-Pr acetate and 60 mL of MeOH for 24 h at 65° and 3 atm of H2.

REFERENCE COUNT: THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L28 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN 2005:490344 CAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 143:43684

TITLE: Process for preparation of optically active

> 3-(4-hydroxyphenyl)propionic acids by reaction of protected 4-hydroxybenzaldehydes and glycolic acid

derivatives to give cinnamates and asymmetric

hydrogenation of the latter.

INVENTOR(S): Yokozawa, Tohru; Shimizu, Hideo; Fujiwara, Takahiro;

Ino, Yasunori

PATENT ASSIGNEE(S): Takasago International Corporation, Japan

SOURCE: PCT Int. Appl., 95 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE

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WO 2005051882
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                               20050609
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                                                                 20041126
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             LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
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                                           EP 2004-819490
     EP 1687250
                               20060809
                                                                   20041126
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     JP 2007512222
                     Τ
                               20070517
                                           JP 2006-520429
                                                                   20041126
     US 20070142472
                         A1
                               20070621
                                           US 2006-578744
                                                                   20060510
                                                               A 20031127
PRIORITY APPLN. INFO.:
                                           JP 2003-398201
                                            WO 2004-JP17998
                                                               W 20041126
OTHER SOURCE(S): CASREACT 143:43684; MARPAT 143:43684
GT
/ Structure 184 in file .gra /
     Title compds. (I; R2 = alkyl; R5-R8 = H, substituent) and salts thereof were
AB
     prepared by reaction of benzaldehydes (II; R1 = protective group; R5-R8 as
     defined above) with R2OCH2CO2R3 (R3 = hydrocarbyl; R2 as defined above),
     hydrolysis of the resulting cinnamate esters to give cinnamic acids, asym.
     hydrogenation, and O-deprotection. Thus, a mixture of 4-
     benzyloxybenzaldehyde, Me methoxyacetate, and NaOMe was refluxed 5 h in MeOH
     to give 80% Me 3-(4-benzyloxyphenyl)-2-methoxyacrylate. This was refluxed 2 h
     with 1N NaOH in MeOH to give 85% 3-(4-benzyloxyphenyl)-2-methoxyacrylic acid
     Na salt. The latter was hydrogenated in MeOH over [Ru(p-cymene)[(S)-dm-
     segphos]]Cl in MeOH at 5 MPa and 60° for 16 h to give Na 3-(4-hydroxyphenyl)-
     2-methoxypropionate in 20% yield and 92.9% enantiomeric excess.
REFERENCE COUNT:
                         4
                               THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
=> s 117 and 'chiral catalyst'
          2028 L17
        129207 'CHIRAL'
            19 'CHIRALS'
        129212 'CHIRAL'
                ('CHIRAL' OR 'CHIRALS')
        827573 'CATALYST'
        823685 'CATALYSTS'
       1060345 'CATALYST'
                 ('CATALYST' OR 'CATALYSTS')
          2138 'CHIRAL CATALYST'
                 ('CHIRAL'(W)'CATALYST')
L29
             0 L17 AND 'CHIRAL CATALYST'
=> s 117 and 'asymmetric react?'
          2028 L17
         76842 'ASYMMETRIC'
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31 'ASYMMETRICS' 76873 'ASYMMETRIC'

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        147848 'ASYM'
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L30
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        171543 'ASYMMETRIC?'
                 ('ASYMMETRIC' OR 'ASYM')
L31
            10 L17 AND 'ASYMMETRIC?'
L31 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
     2007:1187668 CAPLUS Full-text
ΑN
DN
    148:54698
ΤI
    Asymmetric intramolecular alkylation of chiral aromatic imines
     via catalytic C-H bond activation
ΑU
    Watzke, Anja; Wilson, Rebecca M.; O'Malley, Steven J.; Bergman, Robert G.;
    Ellman, Jonathan A.
     Department of Chemistry and Division of Chemical Sciences, Lawrence
CS
     Berkeley National Laboratory, University of California, Berkeley, CA,
     94720, USA
SO
     Synlett (2007), (15), 2383-2389
    CODEN: SYNLES; ISSN: 0936-5214
PΒ
    Georg Thieme Verlag
DT
    Journal
LA
    English
OS
    CASREACT 148:54698
RE.CNT 19
              THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
L31 ANSWER 2 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
AN
    2007:696717 CAPLUS Full-text
DN
    147:95305
ΤI
    Process for the preparation of enantiomer-enriched
     2-alkoxy-3-arylpropionic acids by asymmetric hydrogenation of
     substituted 2-alkoxycinnamic acids
    Woltering, Michael; Bunlaksananusorn, Tanasri; Gerlach, Arne
TN
PA
     Saltigo G.m.b.H., Germany
SO
     Eur. Pat. Appl., 16pp.
     CODEN: EPXXDW
DT
     Patent
LA
    German
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FAN.CNT 1
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            BA, HR, MK, YU
    DE 102005061472
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    US 7429676
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    CN 1986516
                                          CN 2006-10168677
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PRAI DE 2005-102005061472 A
                              20051222
    CASREACT 147:95305; MARPAT 147:95305
             THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 5
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 3 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
L31
    2006:1173484 CAPLUS Full-text
ΑN
DN
    145:489283
ΤI
    N-Acylpiperidines and related compounds as CGRP-antagonists, methods for
    preparing them, pharmaceutical compositions and their use as
    pharmaceutical compositions
    Mueller, Stephan Georg; Rudolf, Klaus; Lustenberger, Philipp; Stenkamp,
IN
    Dirk; Santagostino, Marco; Paleari, Fabio; Schaenzle, Gerhard; Arndt,
    Kirsten; Doods, Henri
PA
    Boehringer Ingelheim International GmbH, Germany
    U.S. Pat. Appl. Publ., 156pp.
SO
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DT
    Patent
    English
LA
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    PATENT NO.
                                                                DATE
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    US 20060252931
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    WO 2005103037
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    WO 2005103037
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                              20060112
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            NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL,
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            ZM, ZW
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EP 1770091
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            BA, HR, MK, YU
PRAI AR 2005-101139
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                         Α
                        А
    WO 2005-EP3094
                               20050323
    WO 2005-EP4104
                        Α
                               20050418
    EP 2005-21283
                        Α
                             20050929
    DE 2004-102004015723 A 20040329
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OS
    MARPAT 145:489283
L31 ANSWER 4 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
    2005:973972 CAPLUS Full-text
AN
    143:422193
DN
ΤI
    Total Synthesis of (+)-Lithospermic Acid by Asymmetric
    Intramolecular Alkylation via Catalytic C-H Bond Activation
    O'Malley, Steven J.; Tan, Kian L.; Watzke, Anja; Bergman, Robert G.;
ΑU
    Ellman, Jonathan A.
    Department of Chemistry, University of California, Berkeley, CA, 94720,
CS
    Journal of the American Chemical Society (2005), 127(39), 13496-13497
SO
    CODEN: JACSAT; ISSN: 0002-7863
    American Chemical Society
PB
DΤ
    Journal
LΑ
    English
OS
    CASREACT 143:422193
RE.CNT 19
             THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 5 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
L31
    2005:490344 CAPLUS Full-text
ΑN
DN
    143:43684
ΤI
    Process for preparation of optically active 3-(4-hydroxyphenyl)propionic
    acids by reaction of protected 4-hydroxybenzaldehydes and glycolic acid
    derivatives to give cinnamates and asymmetric hydrogenation of
    the latter.
    Yokozawa, Tohru; Shimizu, Hideo; Fujiwara, Takahiro; Ino, Yasunori
ΙN
PA
    Takasago International Corporation, Japan
SO
    PCT Int. Appl., 95 pp.
    CODEN: PIXXD2
DT
    Patent
LA
    English
FAN.CNT 1
    PATENT NO.
                       KIND DATE
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                                                                 DATE
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                                          WO 2004-JP17998
    WO 2005051882
                         A1
                              20050609
                                                                 20041126
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            LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
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                        A1 20060809
                                         EP 2004-819490
    EP 1687250
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        R: CH, DE, ES, FR, GB, LI, IE
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	WO	2004-3	JP17998	W	20041126					
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- RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L31 ANSWER 6 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 2003:746329 CAPLUS Full-text
- DN 139:395734
- TI Design and synthesis of small chemical inhibitors containing different scaffolds for lck SH2 domain
- AU Park, See-Hyoung; Kang, Sun-Hee; Lim, Sang-Hyeong; Oh, Hyun-Sik; Lee, Keun-Hyeung
- CS Signal Transduction Laboratory, Mogam Biotechnology Research Institute, Koosung-Myun, Yongin-City, Kyunggi-Do, 449-910, S. Korea
- SO Bioorganic & Medicinal Chemistry Letters (2003), 13(20), 3455-3459 CODEN: BMCLE8; ISSN: 0960-894X
- PB Elsevier Science B.V.
- DT Journal
- LA English
- OS CASREACT 139:395734
- RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L31 ANSWER 7 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
- AN 2003:262905 CAPLUS Full-text
- DN 139:149390
- TI Asymmetric rhodium carbene insertion into the Si-H bond: identification of new dirhodium(II) carboxylate catalysts using parallel synthesis techniques
- AU Buck, Richard T.; Coe, Diane M.; Drysdale, Martin J.; Ferris, Leigh; Haigh, David; Moody, Christopher J.; Pearson, Neil D.; Sanghera, J. Bobby
- CS Department of Chemistry, Loughborough University, Loughborough, LE11 3TU, UK
- SO Tetrahedron: Asymmetry (2003), 14(7), 791-816 CODEN: TASYE3; ISSN: 0957-4166
- PB Elsevier Science Ltd.
- DT Journal
- LA English
- OS CASREACT 139:149390
- RE.CNT 54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L31 ANSWER 8 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN _
- AN 2003:155400 CAPLUS Full-text
- DN 138:338116
- TI Synthesis and Biological and Structural Characterization of the Dual-Acting Peroxisome Proliferator-Activated Receptor α/γ Agonist Ragaglitazar
- AU Ebdrup, Soren; Pettersson, Ingrid; Rasmussen, Hanne B.; Deussen, Heinz-Josef; Jensen, Anette Frost; Mortensen, Steen B.; Fleckner, Jan; Pridal, Lone; Nygaard, Lars; Sauerberg, Per
- CS Novo Nordisk Park, Novo Nordisk A/S, Maalov, 2760, Den.
- SO Journal of Medicinal Chemistry (2003), 46(8), 1306-1317 CODEN: JMCMAR; ISSN: 0022-2623
- PB American Chemical Society
- DT Journal
- LA English

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     CASREACT 138:338116
RE, CNT 72
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              ALL CITATIONS AVAILABLE IN THE RE FORMAT
L31 ANSWER 9 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
     1998:82175 CAPLUS Full-text
     128:167302
DN
OREF 128:32973a,32976a
     A non-enzymic synthesis of (S)-(-)-rosmarinic acid and a study of a
     biomimetic route to (+)-rabdosiin
     Bogucki, David E.; Charlton, James L.
ΑU
     Dep. Chem., Univ. Manitoba, Winnipeg, MB, R3T 2N2, Can.
CS
     Canadian Journal of Chemistry (1997), 75(12), 1783-1794
     CODEN: CJCHAG; ISSN: 0008-4042
    National Research Council of Canada
PΒ
DT
    Journal
LA
    English
    CASREACT 128:167302
OS
RE.CNT 52
              THERE ARE 52 CITED REFERENCES AVAILABLE FOR THIS RECORD
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L31 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2009 ACS on STN
    1989:92028 CAPLUS Full-text
     110:92028
DN
OREF 110:15153a,15156a
     Rabdosiin, a new rosmarinic acid dimer with a lignan skeleton, from
     Rabdosia japonica
     Agata, Isao; Hatano, Tsutomu; Nishibe, Sansei; Okuda, Takuo
ΑU
     Fac. Pharm. Sci., Higashi Nippon Gakuen Univ., Hokkaido, 061-02, Japan
CS
     Chemical & Pharmaceutical Bulletin (1988), 36(8), 3223-5
     CODEN: CPBTAL; ISSN: 0009-2363
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     English
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L8
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L31		10		AND	'ASYMMETRIC?'